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TITLE:

SYSTEM AND METHOD FOR SIMULATING CELLULAR BIOCHEMICAL PATHWAYS

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SYSTEM AND METHOD FOR SIMULATING CELLULAR BIOCHEMICAL PATHWAYS

RELATED APPLICATIONS

Priority is claimed to U.S. Provisional Application No. 60/188,168, filed March 10, 2000, and U.S. Provisional Application No. 60/244,694, filed October 31, 2000, both of which are hereby incorporated by reference in their entirety.

FIELD OF THE INVENTION

The present invention relates to a system and method for examining pathways. In particular, the present invention provides a system and method for examining pathways that underlie cellular functions, specifically signal transduction pathways.

BACKGROUND OF THE INVENTION

DNA sequence analysis and recombinant DNA technology are powerful tools in biologic research. With advances in cellular biology, genetics, and computational methods, a deeper understanding of cell function and disease is emerging. A bulk of current research activity involves efforts to understand the molecular basis of cellular biochemical pathways, *i.e.*, the ordered series of biochemical interactions (typically among proteins) that underlie diverse cellular functions. Greater understanding of these processes will foster more rational and effective design of medicinal therapies.

The recent completion of initial phases of several genome sequencing projects has provided important new resources for understanding cellular biochemical pathways and functions, but more detail is needed to fully understand many cellular functions. For instance, analysis of biochemical pathways, as well as gene and protein functions, are



typically performed with complete knowledge of all the players known to be involved in the relevant cellular biochemical pathways. Thus, the ability to simulate cellular biochemical pathways and probable interaction partners for a protein under investigation would be desirable.

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SUMMARY OF THE INVENTION

One aspect of the invention relates to the prediction of (a) functional properties of a protein, (b) potential interaction partners of the protein, and/or (c) potential target biochemical pathways within which the protein may interact. Thus, according to the invention, the influence of a given stimulus on a biochemical pathway can be assessed.

Another aspect of the invention relates to a system and method for simulating cellular biochemical pathways. The invention integrates the vast information available on cellular biochemical pathways to evaluate and predict the effect of given stimuli on cellular biochemical-pathways. As such, the invention enables investigators working on poorly defined cellular biochemical pathways to simulate the biochemical pathway and predict potential protein interaction partners, in order to gain further insight into possible cellular biochemical pathways in which a target protein may function.

Another aspect of the invention is a system and method for demonstrating the signal cascades that occur in certain cells when certain stimuli are introduced. In an embodiment of the present invention, an inference engine linked to a database of known cellular components and reactions generates the signal cascades.

A further aspect is the incorporation into the system and method of the present invention of DNA sequence analysis of domains, motifs, and sites in new proteins of interest to enable a User to predict the most likely types of upstream and downstream

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proteins (or other biomolecules) with which a new protein might interact and, subsequently, the potential biochemical pathways within which a new protein might act.

This aspect provides new advantages of significantly greater efficiency, confidence, and focus for a User in deciding on potential new avenues of research to pursue.

Another aspect of the system and method of the present invention is the incorporation of data regarding how the primary sequence of functional sites in biomolecules (eg., proteins) effects the specificity and efficacy of physical interactions with binding partners. Further, binding constants, rate equations, and reactant concentrations may be incorporated into the system and method of the present invention, in order to determine reaction events, pathway activities, and cell function outcomes.

The system and method of the present invention may also be used for molecular examination of the relationship between the structure of functional sites and partner interactions and their relation to the effects of molecular interventions by genetic variation, pharmaceutical compounds or toxic substances on the physical interactions of binding partners. Further, the present invention may be used to examine the functional consequences of such molecular interventions to biochemical pathways and cellular events. Examination, with the present invention, of the relationship between sequence variation (molecular genotype) within domains and functional profile within pathways creates new advantages for the design and selection of appropriate pharmaceutical compounds that are unlikely to produce adverse side effects, predicted by the subject genotypic profile.

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or endpoints are introduced. Instead of using pre-generated biochemical pathways, the system and method of the present invention dynamically generate their results using a simulation module that includes an inference engine linked to at least one dynamic database of definitions relating to cellular concepts, components, and reactions.

In one embodiment of the present invention, a method for simulating at least one aspect of a cellular biochemical pathway is provided comprising the steps of: providing information regarding a target cellular environment and a stimulus event; simulating at least one aspect of a cellular biochemical pathway based on the stimulus event and target cellular environment information provided; and textually and/or graphically displaying at least one aspect of a cellular biochemical pathway. A method of the invention can further comprise the steps of predicting target protein functions and/or predicting potential target protein interaction partners.

In an embodiment of the present invention, a method for simulating at least one aspect of a cellular biochemical pathway is provided comprising the steps of: providing information regarding a target cellular environment and an endpoint; simulating at least one aspect of a cellular biochemical pathway based on the endpoint event and the target cellular environment information provided; and textually and/or graphically displaying at least one aspect of a cellular biochemical pathway.

In another aspect of the invention, a system for simulating at least one aspect of a cellular biochemical pathway is provided, comprising: a data input interface; a simulation module; and a display module. Based on cellular environment and input information provided to the data input interface, the simulation module simulates at least one aspect of a cellular biochemical pathway by determining the order of cellular events which occur within the defined cellular environment, and the display module can display textual

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and/or graphical representations of the simulated pathway. Input information may comprise information regarding cellular context, stimuli, knockouts and/or endpoints. The system of the present invention may optionally comprise a prediction module for predicting likely biological outcomes (e.g., apoptosis, lymphocyte activation, etc.), as well as protein interaction partners or gene interaction sites (for transcription factors) based on the simulated pathway(s).

In an embodiment of the present invention, the system and method of the present invention are adapted to be used as an aid in teaching, as educational tools, and/or as a complement to academic textbooks.

In another embodiment, the system and method of the present invention are adapted to be utilized by persons conducting genomic and proteomics research.

BRIEF DESCRIPTION OF THE DRAWINGS

Figure 1A is a block diagram of a system for simulating cellular biochemical pathways, in accordance with one embodiment of the present invention;

Figure 1B is a block diagram of the simulation module of Figure 1A, in accordance with one embodiment of the present invention;

Figures 2A and 2B are a flow chart of a preferred control routine for a forward pathway generation function of the inference engine of Fig. 1B;

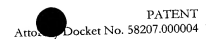
Figure 2C is a flow chart of a preferred control routine for a reverse pathway generation function of the inference engine of Fig. 1B; and

Figures 3 - 43 are examples of various graphical displays that can be generated by the graphical user interface of Fig. 1B.

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DETAILED DESCRIPTION OF THE PREFERRED EMBODIMENTS

Cellular pathways involve molecular physical interactions between elements in series (typically, though not exclusively, proteins) leading to an outcome in a cellular process. Thus, a molecular understanding of these physical interactions, of pathway interconnections, and of pathway architectures would foster a more rational and effective design of pharmaceutical therapies when intervention is needed. Unfortunately, the identity of all the elements and all the interconnections in cellular pathways are not yet established. Gaps exist with respect to many of the elements and their functions. Thus, a major goal of research is to identify all elements of cellular pathways and to understand the molecular functions and interactions of these elements.

Central to this effort is the determination of the DNA sequence of genes that encode these proteins of interest. Examination and comparison of new DNA sequences with known gene DNA sequences stored in public databases routinely provides powerful predictive ability concerning the likely function of a new protein. Deeper examination of sequence data of new proteins can reveal the presence of various functional sites (e.g., domains, motifs, catalytic sites, and sites of biochemical modification), which typically constitute regions of physical interaction (e.g., protein-protein) between biomolecules. Many protein domains, motifs and sites have now been identified and the character of their involvement in diverse molecular interactions are known.

As used herein, the term "cellular biochemical pathway" generally refers to an ordered series of physical interactions between successive cellular elements leading to an outcome (e.g., signal transduction) in a cellular biochemical process.

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Cellular environment information and input information can be provided to the data input interface in any manner known in the art, such as by manual data input through a keyboard or automated data importation.

Protein type information can be derived in any manner known in the art. For instance, a sequence similarity search can be performed, using public or commercial software programs, on a gene or protein sequence of interest. Typical similarity search platforms include the BLAST family of routines available at www.ncbi.nlm.nih.gov/BLAST. Search routines such as BLOCKS, MoST, Pfam, PROSITE, or PROBE that detect conserved protein motifs can then be used if desired. Additional analysis of target protein structure, composition, and function can be done with a variety of web-based platforms.

As used herein, the term "cellular environment" generally refers to the sum total of all the substances and components within a cell under consideration. The provided cellular environment information, according to the present invention, represents at least a portion of the total cellular environment. Such cellular environment information is generally provided as cellular concepts and attributes, which are defined and described in more detail below. Cellular environment information may comprise, but is not limited to, cell type; protein type information, e.g., cell surface trans-membrane receptor; subcellular location; identity of motifs; modification sites; and modification effects, e.g., activation, inhibition, etc. Input information may comprise information regarding stimuli, knockouts and endpoints.

A cellular "concept", as used herein, is an abstraction of anything that can be said to exist in space or occur over time with regard to a cellular environment. For instance, all cellular substances, cellular processes, and cellular components are "concepts". For

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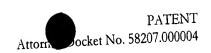
example, in the statement "adenosine binds to an adenosine receptor in a liver cell, which leads to transcription", the concepts are "adenosine", "binds to", "adenosine receptor", "liver cell", "leads to" and "transcription". In grammatical terms, concepts usually represent nouns and verbs.

Figs. 1A and 1B are block diagrams of a system 5 for simulating cellular biochemical pathways, in accordance with one embodiment of the present invention. As shown in Fig. 1A, the system 5 comprises a Simulation Module 10, an Output Module 60, a Report Module 70 and a Database 80. The Simulation Module 10 is in communication with one or more Users 20, an Output Module 60, the Report Module 70, and the Database 80. The Simulation Module 10 contains all the processing logic for the system 5.

The system 5 of the present invention is preferably implemented on a server, which may be or include, for instance, a work station running the Microsoft WindowsTM NTTM, WindowsTM 2000, UNIX, LINUX, XENIX, IBM, AIX, Hewlett-Packard UXTM, NovelTM, Sun Micro Systems SolarisTM, OS/2TM, BeOSTM, Mach, Apache Open StepTM, or other operating system or platform. However, the system 5 of the present invention could also be implemented on a programmed general purpose computer, a special purpose computer, a programmed microprocessor or microcontroller and peripheral integrated circuit elements, an ASIC or other integrated circuit, a hardwired electronic or logic circuit such as a discrete element circuit, a programmable logic device such as a FPGA, PLD, PLA, or PAL, or the like. In general, any device on which a finite state machine capable of implementing the modules and control routines discussed herein can be used to implement the present invention.

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User Interface 12 and an Inference Engine 14 and, optionally, an Editor or Compiler 16. In an embodiment of the present invention, the Graphical User Interface 12 of the Simulation Module 10 may gather input information from a User 20. A User 20 may provide several types of input to the Simulation Module 10 using any data input method known in the art. For example, a User 20 may provide pathway generation parameters to the Simulation Module 10. Further, new data requests may be entered by a User 20 through the Graphical User Interface 12 to the database 80. A User 20 may also input requests for information. Such a request may be entered using a dynamic display. The Simulation Module 10 may also receive input information from the Database 80.

The Inference Engine 14, working with the Database 80, evaluates a sequence of logic statements to determine which cellular events should be triggered based on the cellular environment present at the decision making moment.

As discussed above, the Simulation Module 10 may further comprise an Editor or Compiler 16. The Editor or Compiler 16 may be used by a User 20 to enter new definitions of attributes, concepts and events, to edit existing definitions and/or compile all changes to the Database 80. In a preferred embodiment, a User 20 may open the Database 80 for viewing. In a further embodiment, the User 20 may edit and/or compile attributes so that the results of the edited attributes may be used by the Inference Engine 14 of the Simulation Module 10. The User 20 may also edit and/or compile concepts so that the results of the edited concepts may be used by the Inference Engine 14 of the Simulation Module 10. Additionally, the User 20 may edit and/or

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Engine 14 of the Simulation Module 10. Optionally, a User 20 may be able to save files to the Database 80 using the Editor or Compiler 16 of the Simulation Module 10.

Several outputs may be generated by the Simulation Module 10. For example, requests for data by a User 20 may be sent to the Database 80 by the Inference Engine 14 of the Simulation Module 10. The Simulation Module 10 may generate a static graphical display via the Graphical User Interface 12. The static graphical display may be a display which shows the pathways created with the input information. For example, the static graphical display may be a "step down" or a "mass-action" diagram. This static graphical display or map may then be exported to the Output Module 60 and saved as a separate graphics format file. This graphics file format may be used as a visual aid by a User 20 conducting a presentation.

The Simulation Module 10 may also generate a dynamic graphical display of a "virtual cell" with the pathways that are created with the input information. This dynamic graphical display may be for forward pathway generation. The dynamic graphical display may also be a "virtual" three-dimensional cell. A User 20 may utilize the dynamic graphical display to navigate through the virtual three-dimensional cell. In this manner, the User 20 may look at different substances in the virtual three-dimensional cell as reactions occur. The User 20 may also zoom in and out of the cell and view the cell from different vantage points and perspectives. The dynamic graphical display may also utilize pictures of cells, cell organelles and other pieces of the cell. These pictures may be, for example, from an image created with an electron microscope. In this manner, the dynamic graphical display provides a User 20 with a realistic presentation of the cell.

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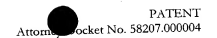
The Simulation Module 10 may additionally generate a written or textual display of the pathway interactions. Such a display may be generated in a display window. The Simulation Module 10 may also communicate information to the Output Module 60 to be used to generate further types of output or results. In a preferred embodiment of the present invention, the Output Module 60 creates a written display of the pathway interactions in a text file. Further, the Simulation Module 10 may communicate information to the Report Module 70 to be used to generate output reports.

The Database 80 preferably stores signal transduction information, which may originate from an operator of the System 5. The information may also originate from the input of a User 20. In a further embodiment, the information may originate from an outside database (not shown) which communicates information to the Database 80. The Database 80 also preferably stores the definitions of specific attributes, concepts, and events. Alternatively, the definitions of specific attributes, concepts, and events may be stored in a separate dynamic definitions database (not shown) that is either a stand-alone database or that is an integrated component of the Simulation Module 10. As an additional alternative, a base definitions database (nor shown) may be incorporated into the Simulation Module 10, and an extended definitions database (not shown) may be compiled and stored externally from the Simulation Module 10.

A User 20 may add new attributes to the Database 80. The Database 80 may also contain information regarding pathologies. This information may comprise signal transduction pathways, as well as different patterns of expression of all the components of the pathways (e.g., protein and DNA level information). For example, a User 20 of the Simulation Module 10 makes a query with a protein differentially expressed. In response, information, not only of the possible associated diseases, but also of the stage of a certain

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disease is preferably provided by the Simulation Module 10. Conversely, a search for a pathology or drug will preferably result in molecular information about that pathology or drug.

The Database 80 may also be coded with information specific to chemical areas which focus on signal transduction within plant or animal cellular environments. For example, the Database 80 may be coded with information specific to pesticides, herbicides, or fertilizers.

As discussed above, a concept is an abstraction of anything that can be said to exist in space or occur over time with regard to a cellular environment. A concept can "inherit" from other concepts, and they can contain other concepts. For example, a User 20 may create a concept of a "protein" and assign certain properties to a "protein" concept. If the User 20 wishes to create an instance of a protein (e.g., "TNF"), the User 20 may define the instance of the protein as a specific type of "protein" and that instance of the protein can inherit all the special properties of a "protein" without having to redefine all these properties a second time. In another example, a User 20 may define all the substances normally contained within mitochondria, and define a "mitochondria" concept that contains all these substance concepts. The User 20 may similarly define other cellular structures with their attending substances and then define a "cell" type to contain all these structures. In this manner, both a hierarchy of types and a hierarchy of structures may be established.

A User 20 may add new concepts to the Database 80. These concepts may be associated with various attributes. In an embodiment of the present invention, these attributes are necessary for efficient processing. For example, the attributes may provide information regarding the shape, color, size or location of a graphic. In a further

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embodiment, the attributes may be informational in nature. For example, the attributes may provide information regarding reference and species. These references may also be tracked by the Simulation Module 10 and exported to the Output Module 60 to facilitate the creation of footnotes, endnotes, articles or reports.

Concepts of the present invention are capable of being inherited from other concepts (e.g., a breast cancer cell inheriting from a generic cell). Concepts may also be capable of containing other concepts (e.g., generic cells containing a nucleus, mitochondria, etc.). Concepts may additionally be capable of excluding other concepts. Further, concepts may be capable of joining other concepts. In a preferred embodiment of the present invention, user input and editing of concept functions is facilitated through a concept wizard. A concept may also be selected by a User 20 to provide details as to the properties of the concept. For example, a User 20 may select a protein to provide the protein sequence with active sites, motifs, signal peptides, etc.

An event is a formal specification of a chemical reaction or process, in terms of (a) the reactants (*i.e.*, what is required for the reaction to occur), (b) the products (*i.e.*, what is produced by the reaction), (c) the inhibitors (*i.e.*, what cannot be present for the reaction to occur), and (d) the context within which the reaction may or may not occur. In this manner, a process defined as a concept (*e.g.*, "gene transcription") is distinguished from one defined as an event, as the event definition requires the pre-conditions and post-conditions of the process to be defined. For example, in the statement, "adenosine binds to an adenosine receptor in a liver cell which leads to transcription", the event is the entire statement, "adenosine" and "adenosine receptor" are the reactants, "gene transcription" is the product and "liver cell" is the context.

A User 20 may add new events to the Database 80 with specific concepts. Events may be associated with various attributes. These various attributes may be necessary for efficient processing (e.g., mobility) or may be informational in nature (e.g., reference and experimental conditions). Events may be capable of requiring the presence of certain concepts in the cellular environment before the event proceeds. Events may also be capable of applying certain concepts to events (e.g., binds, trimerizes, activates, etc.). Further, events may be capable of producing previously defined Events may additionally be capable of being inhibited by user-defined concepts. Events may also be able to specify within which mediums (e.g., cell types) the event may occur and may not occur. For example, the event could specify that the event 10 occurs when contained in a breast cancer cell but not when contained in a liver cell. In a preferred embodiment of the present invention, user input and editing of event functions is facilitated through an event wizard, as will be explained in more detail below.

An attribute is a property of a concept or event. For example, an attribute of a concept may determine what color the Simulation Module 10 should draw the concept if it is represented on a computer screen. In another example, an attribute of an event may provide the Simulation Module 10 with information regarding the length of time the event requires to proceed. In an embodiment of the present invention, a User 20 may add new attributes to the Database 80. The attributes may comprise a decimal, enumeration (e.g., list), integer or text. Further, an attribute comprising a decimal or

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Users 20 simultaneously. In a further embodiment, Users 20 may enter data or input information into the Database 80 and allow the Simulation Module 10 to use that data or input information when generating simulations of various pathways.

In another embodiment of the present invention, a User 20 must supply a predefined user identification code and a corresponding password in order to access the system. The predefined user identification code may be used by the system during the interaction with the database by the User 20. In this embodiment, the User 20 may only be provided access to certain predefined areas of the Database 80 which correspond to the access level assigned to the User 20. The predefined user identification code may also be assigned to a group identification code that corresponds to another predefined access level. In addition, to access to certain predefined areas of the Database 80, the user or group identification code may also provide access to a second database (not shown). In this manner, the User 20 may have access to at least one database of information with the amount of access corresponding to a user or group identification code.

The Simulation Module 10 may generate a forward cell pathway. The forward cell pathway may be generated by prompting a User 20 to specify (a) the cell type where the simulation will be conducted and/or (b) the stimulus/stimuli (i.e., initiating event) for the event. The User 20 may also exclude certain events from or designate reaction endpoints for the forward cell pathway. These user inputs may be facilitated by an input

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compare certain concepts of two or more different concepts to determine if an event should proceed.

As illustrated in Fig. 1A, the Simulation Module 10 utilizes information provided by the Database 80 and by Users 20. The Inference Engine 14 of the Simulation Module 10 then processes the information to predict pathways. In one embodiment of the present invention, the Inference Engine 14 may predict a biochemical signal transduction pathway, a small molecule metabolic pathway, a detoxification enzyme pathway (e.g., a P450 enzyme-mediated biotransformation to various metabolites), toxicology, acute phase reactions, or complement cascades (e.g., classical, alternate, and MBL). In another preferred embodiment of the present invention, the system of the present invention may generate diagrams, assist in interpreting the outcome of gene expression (i.e., functional genomics), and/or facilitate drug target identification or validation.

In another preferred embodiment, the system and method of the present invention may interface with microarrays to predict changes in the activity of biochemical pathways in response to diverse conditions. In this embodiment, the activities of biochemical pathways vary in response to altered conditions. For example, a common response to variation in the activity of biochemical pathways is a change in the expression of many target genes. In another example, variation in the amount of mRNA and the resultant effect on proteins made from the target genes contribute to changes in cell function.

In a further preferred embodiment, microarray chips provide an efficient means to rapidly survey quantitative changes in the expression of a large set of genes that result

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proteins translated from the mRNAs expressed by the target genes in a given cell type under differing conditions could be rapidly detected on a chip. Further, a change in the amount of protein made by a gene would likely change the activity of the biochemical pathway in which the protein product functioned. In this manner, the information obtained from microarray chips may be imported to the Simulation Module 10 to evaluate how changes in the expression of large gene sets might change the activity of diverse biochemical pathways in response to varied conditions. Further, the Simulation Module 10 may be used to examine responses to many conditions such as response to presentation of pharmaceutical or toxic substances or the character of cell function under pathologic versus normal conditions (e.g., in a B lymphoma cell line verses a normal B cell).

Transcriptional (microarray) and translational (protein chip) data may be directly uploaded into the Database 80. The Inference Engine 14 of the Simulation Module 10 may infer potential signal transduction pathways at the global level and model the pathways. In a further embodiment of the present invention, the Simulation Module 10 contains sufficient data to not only reconstruct normal cellular pathways but also pathways associated with disease states. Expression data on gene products in biochemical pathways may also be input into the pathways generated by the Simulation Module 10. The pathways and interactions of drugs may be subsequently modeled and additional simulations may be generated regarding drug action on normal and diseased cells and/or organs. The Simulation Module 10 may also simulate signal transduction pathways at the cell level to provide output information regarding the role and physiological importance of new gene products or gene products with altered expression.

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as potential drug targets. In this embodiment, the list of possible or potential drug targets may also be expanded and reprocessed by the Simulation Module 10. In another preferred embodiment, the present invention simulates signal transduction pathways which are subsequently used to examine the therapeutic value and possible toxicities of drug candidates.

The system and method of the present invention may further comprise a high throughput screening system (not shown) or other automated assay testing device (not shown). This automated assay testing device would provide a User 20 with an automated method to perform simulated pathways in a laboratory.

The system and method of the present invention may also incorporate the cojoining of multiple cell types to form multi-functional tissues, the co-joining of tissues to form organs, and the co-joining of multiple organs to form organ systems. In this manner, the Inference Engine 14 of the Simulation Module 10 may predict whole body responses, via signal transduction modeling, to individual and multiple stimuli.

The system and method of the present invention may be used as an aid in teaching or educational tools, or as complement to academic text books. Academic institutions offering course work in the life sciences could benefit from using the system and method of the present invention as a powerful teaching and research tool. Students often comprehend difficult concepts better when they are presented in an interactive and visual manner. The system and method of the present invention provides static and dynamic pathway displays that can be used by educators to teach students about the complexities of the intracellular environment, including the interrelationships of multiple components and pathways. In addition, the present inventions' ability to incorporate new concepts and events can facilitate learning about the maturation of cells to fully

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differentiated states, the progression of disease processes like cancer in cells, and the interaction of pharmaceutical products with each other and cellular components.

In addition, the system and method of the present invention can easily be adapted to incorporate the specific concepts and events associated with other types of cellular-based organisms. Thus, for example, the system and method of the present invention can be used in the agricultural industry. The U.S. Department of Agricultural has initiated several national programs related to plant and animal production, product value, and safety through its Agricultural Research Service (ARS). ARS researchers in many of these national programs could benefit from using the system and method of the present invention, because their work also focuses on achieving a better understanding of intracellular interactions.

In an embodiment of the present invention, a User 20 may simulate assays using the systems and methods of the present invention. In a further embodiment of the present invention, the Database 80 and Inference Engine 14 may be accessed via an internet website. A User 20 may generate forward and reverse pathways through the internet website. Should a User 20 wish to physically run the simulated assays, the User 20 may obtain the necessary tools via hyperlinks corresponding to materials utilized in the simulation. For example, a User 20 may select a hyperlink corresponding to a concept or event and information may be presented regarding purchase information for assay kits or reagents. In a further embodiment, a User 20 may select a hyperlink corresponding to materials utilized in the simulation and the User 20 would be presented

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Fig. 2A is a flowchart of a preferred control routine for a forward pathway generation function of the Inference Engine 14 of Fig. 1B. The control routine begins at step 18, where the Inference Engine 14 accesses the Database 80 to create the cellular environment and pathway data structures. For example, the Inference Engine 14 may retrieve values and indicia corresponding to data implicated by the environment and pathway.

Control then continues to step 21, where those substances present in the cellular context are added to the environment. These substances may be determined based upon information stored in the Database 80. Then, at step 22, the substances selected by a User 20 as stimuli are added to the cellular environment and pathway data structures. Control then continues to step 24.

At step 24, any concepts that a User 20 has specified as "knockouts" are removed from the cellular environment and pathway data structures. Control then continues to step 26, where all substances in the cellular environment that are determined to be currently available to the pathway are marked.

Next, at step 28, the first reaction defined in the database is designated as the current reaction. Control then continues to step 30, where it is determined whether there is a substance present in the cellular environment that would inhibit the current reaction. If there is, control jumps to step 42, where the reaction is discarded and control continues to step 46. Otherwise, control continues to step 32.

At step 32, it is determined whether all the reactants of the current reaction are available in the cellular environment. If not all the reactants are available, control jumps to step 42. Otherwise, control continues to step 34.

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At step 34, it is determined whether any additional user-defined tests of the reaction fail. If so, control jumps to step 42. Otherwise, control continues to step 36, where the Inference Engine 14 adds the reaction to the sequence of events that make up one, or more, of each cellular pathway.

Next, at step 38, the Inference Engine 14 adds the products of the reaction to the environment and uses the reaction duration to show relative process time of the products. Control then continues to step 46, where the Inference Engine 14 determines if all the reactions have been tested. If not, control jumps to step 44. Otherwise, control continues to step 50.

At step 50, the Inference Engine 14 identifies whether any endpoints defined by the User 20 have been reached. If so, the control routine ends. Otherwise, control continues to step 52.

At step 52, the Inference Engine 14 determines whether all the substances in the cellular environment have been activated. If so, the control routine ends. Otherwise, control jumps to step 26.

At step 44, the next reaction defined in the Database 80 is designated as the current reaction. Control then jumps to step 28.

Fig. 2B is a flowchart of a preferred control routine for a reverse pathway generation function of the Inference Engine 14 of Fig. 1B. The method begins at step 54, where the Inference Engine 14 accesses the Database 80 to add to the environment those substances selected by the user as endpoints. Control then continues to step 56, where the Inference Engine 14 identifies each underived product in the products.

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Next, at step 58, for each underived product identified in step 56, the Inference Engine 14 determines whether the product is a stimulus. If so, control returns to step 56. Otherwise, control continues to step 62.

At step 62, all events in the particular cellular environment which may produce the identified stimulus product are added to the product's list of producers. Control then continues to step 64, where, for each producer, a new product is generated for each of the corresponding reactants. Each of the current product's producers is subsequently added to the new product's list of consumers. Control then continues to step 66.

At step 66, it is determined whether there are any remaining underived products. If so, control jumps back to step 56. Otherwise, control continues to step 68.

At step 68, a link between each producer and each consumer is established for each product. Control then continues to step 69, where all events are added to the pathway. The control routine then ends.

In one embodiment of the invention, the definitions are stored in the Database 80 in a binary format, an editable textual format, or a combination of both. The editable textual format preferably comprises a descriptive computer language called Signal Transduction Language ("STL"), which is one aspect of the present invention. The definitions may be generated using STL, and/or compiled into a binary format in any manner known in the art. In a preferred embodiment of the invention, definitions of attributes, concepts, and events are created, and cellular biochemical pathways are simulated using a set of graphics-based forms (*i.e.*, instructive, step-by-step screens) generated by the Simulation Module 10. In an alternate preferred embodiment, a set of graphics-based forms are generated by the Output Module 60. Alternatively, definitions and simulations may be created through a direct, text-based interface using an STL shell.

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As described above, in one embodiment of the present invention, the Graphical User Interface 12 may be implemented with graphics-based forms. In this embodiment, upon initiating the Simulation Module 10, a User 20 may be presented with a main menu 100, an example of which is shown in Fig. 3. To perform an action, the User 20 may select one of a plurality of user action buttons 110a-112g. Each of the user action buttons 110a-110g may be associated with one of a plurality of Simulation Module functions. In the main menu 100 shown in Fig. 3, user action buttons 110a-112g may be associated with the Simulation Module functions Define an Attribute; Define a Substance or Process; Define a Reaction; Use STL Editor/Compiler; Generate Pathway; Generate Reverse Pathway; and Exit Program, respectively.

Selecting the Define an Attribute button 110a initiates an Attribute Forms series, which is a series of screens that allows the User 20 to define a new attribute. Selecting the Define a Substance or Process button 110b initiates a Concept Forms series, which is a series of screens that allows the User 20 to define a new concept. Selecting the Define a Reaction button 110c initiates an Event Forms series, which is a series of screens that allows the User 20 to define a new event. Selecting the Use STL Editor/Compiler button 110d initiates an STL shell, which allows the User 20 to directly edit and compile the textual definitions. Selecting the Generate Pathway button 110e or Generate Reverse Pathway button 110f initiates a Pathway Forms series, which is a series of screens that allows the User 20 to initiate the Inference Engine 14 and simulate at least one aspect of a cellular biochemical pathway. Finally, selecting the Exit this Program button 110g shuts down the Simulation Module.

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Attribute Forms series

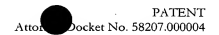
When a User 20 initiates the Attribute Forms series from the main menu 100 by selecting the Define an Attribute button 110a, an Attribute Name screen 200, an example of which is shown in Fig. 4A, may be displayed. As shown in Fig. 4A, a User 20 may have the option of entering an attribute name in field 201 and then selecting the Next button 205 to proceed to the next step in the Attribute Forms series. Alternatively, a User 20 may select the Cancel button 215 to exit the Attribute Forms series. It should be noted that user input in field 201 may determine the step that comes "next" in the Attribute Forms series.

Entering an attribute name in field 201 and clicking the Next button 205 may initiate a Select Attribute Type screen 220, an example of which is shown in Fig. 4B. As shown in Fig. 4B, the Select Attribute Type screen 220 may allow a User 20 to select one of four types of attributes: Decimal 221a; Integer 221b; Text 221c; or Enumerated 221d. A User 20 may select one of the attribute types and then select the Next button 225 to proceed to the next step of the Attribute Forms series. Alternatively, a User 20 may select the Back button 230 to return to the previous screen or the Cancel button 235 to exit the Attribute Forms series.

If a User 20 selects Decimal 221a or Integer 221b, followed by the Next button 225, the Enter Attribute Limits screen 240, an example of which is shown in Fig. 4C, may be displayed. As shown in Fig. 4C, a User 20 may enter numeric attribute values, through the Enter Attribute Limits screen 240, that represent an upper limit 241b and a lower limit 241a for this attribute. Each of the numeric attribute limits may be made inclusive by selecting a respective Inclusive indicator 242a and 242b. A User 20 may select the Next button 245 to proceed to the next step of the Attribute Forms series, the

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Back button **250** to return to the previous screen, or the Cancel button **255** to exit the Attribute Forms series.

Referring back to Fig. 4B, if a User 20 selects Enumerated 221d from the Select Attribute Type screen 220, followed by the Next button 225, the Enumerated Enter Attribute Limits screen 260, an example of which is shown in Fig. 4D, may be displayed. As shown in Fig. 4D, a User 20 may enter all possible values of an enumerated list. For example, the values represented by a "cellular location" type of attribute may be entered by typing the text "membrane bound" into the Value field 261, clicking the Add Value button 262, then repeating the process for any other values represented by the attribute, e.g., "extracellular", "nuclear", etc. Values that have been added may appear in the Allowed Values list 264. From the Enumerated Enter Attribute Limits screen 260, a User 20 may also remove a value from the Allowed Values list 264 by selecting the value from the Allowed Values list 264 and then selecting the Remove Value button 263. A User 20 may select the Next button 265 to proceed to the next step of the Attribute Forms series, the Back button 270 to return to the previous screen, or the Cancel button 275 to exit the Attribute Forms series.

Upon selecting the Next button 265 from the Enumerated Enter Attribute Limits screen 260 (Fig. 4D), the Next button 245 from the Numeric Enter Attribute Limits screen 240 (Fig. 4C), or Text 221c from the Select Attribute Type screen 220 (Fig. 4B), the Attribute Definition Complete screen 280, an example of which is shown in Fig. 4E, may be displayed. As shown in Fig. 4E, the Attribute Definition Complete screen 280 may show, in field 281, the STL code that was produced from user selections within the Attribute Forms series. In the example shown in Fig. 4E, "Sample Attribute" was entered as an attribute name in field 201 (Fig. 4A), the type selected was Decimal 221a

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(Fig. 4B), a value "1" was entered as the lower limit 241a, and a value of "2" was entered as the upper limit 241b (with the inclusive indicators 241a and 242b checked). As shown in Fig. 4E, from the Attribute Definition Complete screen 280, a User 20 may select the Finish button 285 to enter the attribute definition into at least one dynamic database of definitions. A User 20 may optionally select the Back button 290 to return to the previous screen, or may select the Cancel button 295 to exit the Attribute Forms series.

Concept Forms series

When a User 20 initiates the Concept Forms series from the main menu 100 by selecting the Define a Substance or Process button 110b (Fig. 3), a Concept Name screen 300, an example of which is shown in Fig. 5A, may be displayed. As shown in Fig. 5A, a User 20 has the option of entering a concept name in field 301, and then selecting the Next button 305 to proceed to the next step of the Concept Forms series. Alternatively, a User 20 may select the Cancel button 315 to exit the Concept Forms series. It should be noted that user input in field 301 may determine the subsequent step in the Concept Forms series.

Entering a concept name in field 301 and clicking the Next button 305 may initiate the Select a Base Concept screen 320, an example of which is shown in Fig. 5A. The Select a Base Concept screen 320 may allow a User 20 to select a base concept from which a newly named concept will inherit attributes. This is especially useful in defining reactants and products of reactions. For example, in the reaction "A phosphorylates B", the User 20 may define "A" and "B" using "protein" as the base concept. The concept "phosphorylated B" can then be defined using "B" as the base concept. A concept does not require a base concept. However, a User 20 may select one or more base concepts and then select the Next button 325 to proceed to the next step of the Concept Forms

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series. Alternatively, a User 20 may select the Back button 330 to return to the previous screen or the Cancel button 335 to exit the Concept Forms series.

Upon selecting the Next button 325, the Select Any Contained Concepts screen 340 may be displayed, an example of which is shown in Fig. 5C. The Select Any Contained Concepts screen 340 allows a User 20 to select one or more concepts which the concept named in the Select a Base Concept screen 320 (Fig. 5B) is to contain. This feature may be used to create a "cell" or some other general cellular environment. As defined and described in more detail below, the concepts that are contained in a given cellular environment are assumed to be "available" to any biochemical pathway that involves the contained concepts.

Referring to Fig. 5C, if more than one of a selected contained concept is present within a given cellular environment, upon selecting the concept, the number present may be entered in Quantity field 341. For example, a cell which includes a TNFR receptor complex may be defined as containing three TNFR concepts. A User 20 may select one or more contained concepts, enter the number of individual concepts contained in Quantity field 341, and then select the Next button 345 to proceed to the next step of the Concept Forms series. Alternatively, a User 20 may select the Back button 350 to return to the previous screen or the Cancel button 355 to exit the Concept Forms series.

Upon selecting the Next button 345 (Fig. 5C), the Select any Excluded Concepts

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button 365 to proceed to the next step of the Concept Forms series, the Back button 370 to return to the previous screen, or the Cancel button 375 to exit the Concept Forms series.

Upon selecting the Next button 365, if the newly named concept includes the "substance" attribute (which is true for the "substance" concept or any of its inherited concepts, like "protein"), the Customize Concept Appearance screen 380 may be displayed, an example of which is shown in Fig. 5E. The appearance of the concept's graphical presentation may be customized in the Customize Concept Appearance screen 380. The shape of the concept graphic may be selected from the Shape Box 382. The relative size of the concept graphic can be selected in the Relative Size Box 383, or the up and down buttons 383a and 383b may be used to increase or decrease the relative size incrementally. The color of the concept graphic may be selected by selecting the Color button 384. View Box 381 may show the concept as it will appear, and may be updated after every change. A User 20 may select the Next button 385 to proceed to the next step of the Concept Forms series, the Back button 390 to return to the previous screen, or the Cancel button 395 to exit the Concept Forms series.

When the Color button 384 is selected, a standard Windows® color palette screen 384a may be displayed, an example of which is shown in Fig. 5F. A basic color may be selected from the Basic Colors palette 384b, or a custom color may be defined in the Custom Colors palette 384c by manipulating parameters of Color Diagram 384d and selecting the Add button 384e. Once the color is selected, the OK button 384f may be selected to return to the Customize Concept Appearance screen 380 of the Concept Forms series. Alternatively, the Cancel button 384g may be selected to return to the Customize Concepts Appearance screen 380 without defining a color.

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Upon selecting the Next button 385, the Assign Concept Attributes screen 3000 may be displayed, an example of which is shown in Fig. 5G. Attributes associated with the newly named concept can be selected from the Assign Concept Attributes screen 3000. If the newly named concept inherits concepts from a base concept, the inherited attributes may already be selected. Again, a User 20 may select the Next button 3025 to proceed to the next step of the Concept Forms series, the Back button 3050 to return to the previous screen, or the Cancel button 3075 to exit the Concept Forms series.

Upon selecting the Next button 3025, the Assign Attribute Values screen 3100 may be displayed, an example of which is shown in Fig. 5H. An attribute value may be defined by selecting an attribute from the Attribute Box 3110, and entering a desired value in the Value Box 3120. For example, selecting "comments" in the Attribute Box 3110 may allow textual comments to be entered in the Value Box 3120. If the attribute is an enumerated list, a list of allowed values for that attribute may be displayed in the Value Box 3120 for selection. A User 20 may select the Next button 3125 to proceed to the next step of the Concept Forms series, the Back button 3150 to return to the previous screen, or the Cancel button 3175 to exit the Concept Forms series.

Upon selecting the Next button 3125, the Concept Definition Complete screen 3200 may be displayed, an example of which is shown in Fig. 5I. The Concept Definition Complete screen 3200 may show the STL code that may be produced from the User 20 selections. In the example shown in Fig. 5I, the following selections were made: "Sample Concept" was entered as the name (using the screen shown in Fig. 5A); "protein" was selected as the base concept (using the screen shown in Fig. 5B); "activators", "comments", "inhibitors", "location", "other databases" and "references" were selected as attributes (using the screen shown in Fig. 5G); values were assigned to

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"graphiccolor", "graphicshape", and "graphicsize" (using the screen shown in Fig. 5E); and the text string "this is a sample concept" was assigned to "comments" (using the screen shown in Fig. 5H). Once the concept definition is complete, the STL code may be compiled into the definitions database by selecting the Finish button 3225. Alternatively, the Back button 3250 may be selected to return to the previous screen, or the Cancel button 3275 can be selected to exit the Concept Forms series without compiling the concept definition into the definitions database.

Event Forms series

When a User 20 selects Define a Reaction 110c from the main menu 100 (Fig. 3), the Event Forms series may initiate to display an Enter Event Description screen 400, an example of which is shown in Fig. 6A. The Event Forms series may be used to define a new chemical reaction or relationship. As shown in Fig. 6A, a User 20 may have the option of entering an event description in field 401, and then selecting the Next button 405 to proceed to the next step of the Event Forms series. The event description is used for display only and is not parsed or translated by the Event Forms series. Alternatively, a User 20 may select the Cancel button 415 to exit the Event Forms series. It should be noted that user input in field 401 may determine the next step in the Event Forms series.

Upon selecting the Next button 405, the Select Reactants screen 420 may be displayed, an example of which is shown in Fig. 6B. One or more reactants for the event may be selected from the Selection Box 421. According to the present invention, reactants may be any concept required for a cellular reaction or biochemical pathway to proceed. In the case of a molecule binding to a receptor, for example, both the stimulus molecule and the receptor molecule are reactants and must be selected. If more than one of a given concept must be present for a reaction to proceed, that number may be

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entered in the Quantity Box 422 after the concept is selected from the Selection Box 421.

A User 20 may select the Next button 425 to proceed to the next step of the Event Forms series, the Back button 430 to return to the previous screen, or the Cancel button 435 to exit the Event Forms series.

Upon selecting the required reactants in Selected Box 421, optionally entering a reactant quantity in Quantity Box 422, and selecting the Next button 425, the Select Products screen 440 may be displayed, an example of which is shown in Fig. 6C. One or more products from the reaction may be selected from the Selection Box 441. The products are concepts produced by the process of the reaction. According to the present invention, processes (e.g., apoptosis and gene transcription) may be products. For example, when the process of the reaction is a molecule binding to a receptor, the product might be a bound or activated receptor. If the reaction produces more than one instance of a concept, that number can be entered into the Quantity Box 442 after the product is selected. A User 20 may select the Next button 445 to proceed to the next step of the Event Forms series, the Back button 450 to return to the previous screen, or the Cancel button 455 to exit the Event Forms series.

Selecting the Next button 445 may initiate the Select Inhibitors screen 460, an example of which is shown in Fig. 6D. One or more inhibitors for the reaction may optionally be selected from the Selection Box 461. Inhibitors may be concepts whose presence in the cellular environment means that the reaction cannot proceed. If the reaction is inhibited by more than one instance of a concept, the quantity of concepts may be entered into the Quantity Box 462 after the inhibitor is selected in the Selection Box 461. A User 20 may select the Next button 465 to proceed to the next step of the

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Event Forms series, the Back button 470 to return to the previous screen, or the Cancel button 475 to exit the Event Forms series.

Upon selecting the Next button 465, the Enter Reaction Structure screen 480 may be displayed, an example of which is shown in Fig. 6E. The structure of the reaction, i.e., how the reaction components should interact, may optionally be defined from this screen. Reaction structure definitions are not required for the simulation of the reaction or cellular biochemical pathways in general, but instead may be utilized, for example, to define how the reactions will be textually and/or graphically displayed. To define a reaction structure, a process may be selected from the Process Box 481, the concept from which the process is applied may be selected from the Applied From Box 482, and the concept to which the process applies may be selected from the Applied To Box 483. For example, when the process of the reaction is a molecule "A" binding to a receptor "B", a User 20 may define the relationship "A binds to B" by selecting "A" in the Applied From Box 482, "binds to" in the Process Box 481, and "B" in the Applies To Box 483. The Add button 484a may then be selected to add the reaction structure to the Descriptions List Box 480a. Alternatively, a reaction structure definition may be removed from the Descriptions List Box 480a by selecting the definition from the Descriptions List Box 480a and clicking the Remove button 484b. A User 20 may select the Next button 485 to proceed to the next step of the Event Forms series, the Back button 490 to return to the previous screen, or the Cancel button 495 to exit the Event Forms series.

Upon selecting the Next button 485 from the Enter Reaction Structure screen 480 (Fig. 6E), the Enter Event Attributes screen 4000 may be displayed, an example of which is shown in Fig. 6F. Certain event attributes may optionally be assigned from this

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screen if relevant. For example, the event duration (i.e., the amount of time required for the event to proceed) may be entered in the Event Duration Box 4010. Reactant mobility characteristics and post-reaction presence may also be assigned by selecting the reactant from the Reactants List Box 4020 and checking Mobility Box 4030 and/or Post-Reaction Presence Box 4040. Preferably, checking the Mobility Box 4030 communicates that the corresponding reactant will move toward another reactant or reactants when the reaction is simulated. Likewise, checking the Post-Reaction Presence Box 4040 communicates that the corresponding reactant will be present after the reaction has occurred. A User 20 may select the Next button 4050 to proceed to the next step of the Event Forms series, the Back button 4060 to return to the previous screen, or the Cancel button 4070 to exit the Event Forms series.

Selecting the Next button 4050 from the Enter Event Attributes screen 4000 (Fig. 6F) may initiate the Enter Event Contexts screen 4100, an example of which is shown in Fig. 6G. Cellular environments may be specified from the Enter Event Contexts screen 4100. Applicable cellular environments may comprise: (1) selecting, in Present Box 4110, cell types in which the reaction is present; or (2) selecting, in the Absent Box 4120, cell types in which the reaction is not present. If no cell types are selected in Present Box 4110 or Absent Box 4120, the reaction will be applied to all cellular environments. A User 20 may select the Next button 4130 to proceed to the next step of the Event Forms series, the Back button 4140 to return to the previous screen, or the Cancel button 4150 to exit the Event Forms series.

Upon selecting the Next button 4130, the Event Definition Complete screen 4200 may be displayed, an example of which is shown in Fig. 6H. Preferably, the Event Definition Complete screen 4200 shows the STL code that is produced from the user

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selections throughout the Event Forms series. Referring to Fig. 6H, for example, "adenosine binds to adenosine receptor" was entered as the description (using the screen shown in Fig. 6A), "adenosine" and "A2aR" were selected as reactants (using the screen shown in Fig. 6B), and "bound A2aR" was selected as the product (using the screen shown in Fig. 6C). A reaction description was entered with process "binds to" being applied from "adenosine" to "A2aR" (using the screen shown in Fig. 6E), "generic cell" was selected as the event context (using the screen shown in Fig. 6G) and the Mobility Box 4030 and Post-Reaction Presence Box 4040 were checked (using the screen shown in Fig. 6F). When a User 20 selects the Finish Button 4210, the STL code is preferably compiled into the definitions database. The Back button 4220 may be selected to return to the previous screen or the Cancel button 4230 may be selected to exit the Event Forms series without compiling the concept definition into the definitions database.

STL Editor/Compiler

When Use STL Editor/Compiler 110d is selected from the main menu 100 (Fig. 3), the STL Editor/Compiler screen 500 may be displayed, an example of which is shown in Fig. 7. The STL Editor/Compiler screen 500 may be used to enter new definitions of attributes, concepts and events, to edit existing definitions, and compile changes to the definitions database. The STL Editor/Compiler may also be used to open a script file, enter new definitions or edit existing ones, compile the script to place the changes into the definitions database, and save the script changes. The script files represent an editable, user-readable representation of the definitions contained in the definitions database.

As shown in Fig. 7, in one embodiment of the invention, the menu options available from the STL Editor/Compiler may include File menu 510, Edit menu 520, and

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Tools menu 530. Action options within the File menu 510 may include New, Open, Save, Save As, and Exit using standard Windows dialogs or any other mechanism known in the art. Action options within the Edit menu 520 may include Cut, Copy, and Paste functions. Such Edit menu 520 options may be accomplished using any mechanism known in the art. Action options within the Tools menu 530 include Compile Script and Compile Incrementally functions.

The Compile Script function may compile the current script into the definitions database and replaces the previous entries in the definitions database. The Compile Incrementally function may compile the current script into the definitions database without replacing the previous entries in the definitions database. In a preferred embodiment, if script errors are detected, an error message may be displayed specifying the error that was found, and highlighting the area in the script where the error occurred. In this embodiment, a User 20 may rebuild the complete definitions database by executing the Compile Script function to destroy the old database and add in basic definitions. Alternatively, a User 20 may execute the Compile Incrementally function to add user-defined definitions. A more detailed description of STL is printed below.

Pathway Forms series

The Pathway Forms series may be used to specify the stimulus and context of a potential pathway. This process may start the Inference Engine 14 and generate all possible pathways, applying all known reactions, until no new intermediate products are produced. The program may then display three views of the pathway: textual, static and dynamic. When a User 20 initiates the Pathway Forms series from the main menu 100 by selecting the Generate pathway button 110e (Fig. 3), a Choose a Context for the Pathway screen 600 may be displayed, an example of which is shown in Fig. 8A. As

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shown in Fig. 8A, the Next button 605 will take you to the next step of the Pathway Forms series, the Back button 610 will take you to the previous step, and the Cancel button 615 will exit the Pathway Forms series.

As shown in Fig. 8A, the User 20 may select a cellular context for a pathway. This may be a type of cell. Selecting a Next button 605 may bring the User 20 to a Choose a Pathway Stimulus screen 700, an example of which is shown in Fig. 8B. The Choose a Pathway Stimulus screen 700 may allow the User 20 to choose one or more stimulus concepts. Stimulus concepts may be concepts that may be introduced to the cellular context selected in the Choose a Context for the Pathway screen 600 shown in Fig. 8A. The User 20 may select concepts from the List Box 701 to serve as stimuli. If more than one instance of a concept is required to generate a desired result, that number may be entered in the Quantity Box 702.

Clicking the Next button **705** may bring the User **20** to the Choose Pathway Intermediate(s) to Knock Out screen **800**, an example of which is shown in Fig. 8C. If a User **20** wishes to test how a pathway is influenced by the absence of certain concepts, these concepts may be selected from in the List Box **801**. The selected concepts will be removed from the cellular context. Preferably, however, the selected concepts are only removed from the immediate corresponding pathway generation. Preferably, the selected concepts are not permanently removed from the pathway generation.

Clicking the Next button 805 brings the User 20 to the Choose any Pathway Endpoints screen 900, an example of which is shown in Fig. 8D. If the User 20 wishes the pathway to stop when a certain concept has been generated, such pathway endpoints may be selected in Selection Box 901. This is useful in determining if a pathway intermediate is produced in complex pathways.

Clicking the Next button 905 brings the User 20 to the Pathway Definition Complete screen 1000, an example of which is shown in Fig. 8E. The User 20 may click the Finish button 1005 to initiate the display module, which will be explained in more detail below.

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Display Module

A pathway may be displayed in a textual form, as shown in Fig. 9. Fig. 9 illustrates an example of a Textual Pathway window 1100, which shows each reaction/event in the order that it was triggered. If there are more events than will fit on the Textual Pathway window 1100, a scroll bar may appear to the right to scroll the output. The event(s) that is currently being displayed in the dynamic pathway may be highlighted. The Textual Pathway window 1100 may be sized independently, and may be closed without affecting program operation. In a further embodiment, the Textual Pathway window 1100 may be ordered by pathway with a line between each pathway. In this embodiment, the User 20 has an option to order the events based upon paths or to order the events based upon a time stamp assigned to the path. Preferably, the displayed pathway may be highlighted as it occurs.

The pathway may also be displayed in a static, graphical form, as shown in Fig. 10A. Fig. 10A illustrates an example of a Static Pathway window 1200, which shows a schematic diagram of the concepts involved and the reactions that occur. If the diagram extends beyond the confines of the Static Pathway window 1200, scroll bars may appear on the bottom and right. The Static Pathway window 1200 may be sized independently, and may be closed without affecting program operation.

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In one embodiment of the present invention, a pop-up menu of attributes may appear when the user clicks on a left mouse button on a concept. From the menu of attributes the User 20 may choose from attributes such as "references" or "comments". Choosing an attribute may activate a read-only version of the concept editor. The event editor may also be activated by clicking on the event arrow corresponding to an event.

In one embodiment of the present invention, if the User 20 clicks a left mouse button outside of a concept, a pop-up menu may appear allowing the User 20 to "Cancel" the menu or "Print" the diagram. If the User 20 chooses to print, a Print window 1300 may appear, an example of which is shown in Fig. 10B. The User 20 may make any changes desired, then click the OK button 1305 to print the diagram.

The pathway may also be displayed as a dynamic animation, an example of which is shown in Fig. 11. Fig. 11 illustrates a Dynamic Pathway window 1400, which shows an animation of the pathway against the backdrop of a "standard" cell. A shape represents each concept. The names of each concept are preferably displayed on the sides, with lines preferably drawn from the names to the concept shapes. The current elapsed time of the pathway may be displayed on the bottom right. The Dynamic Pathway window 1400 may be sized independently, and may be closed without affecting program operation.

If the pathway proceeds towards proliferation, apoptosis, differentiation, or other pre-determined events, the display may display the event name on the static display. Gene transcriptions may be represented by a standard symbol to the left of the nucleus, displaying the name of the gene or "gene" id, if it is not known.

As each event is reached in the animation, it may be highlighted in the Textual Pathway window 1100 (Fig. 9).

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In one embodiment of the present invention, if the User 20 clicks the left mouse button on a concept, a pop-up menu of attributes may appear (such as "references", "comments") that the User 20 can choose from. Choosing an attribute may brings up a hyperlink window (Fig. 12). Choosing an attribute may also activate a read-only version of the concept editor. The event editor may also be activated by clicking on the event arrow corresponding to an event.

In one embodiment of the present invention, if the User 20 clicks the left mouse button outside of a concept, a pop-up menu may appear that allows the User 20 to "Cancel" the menu, "Run" the animation, "Stop" the animation, "Restart" the animation from the beginning, or change the "Speed" of the animation. The animation may be stopped and started using "Run" and "Stop", and choosing "Speed" preferably brings up another menu of speeds from 1 (slowest) to 10 (fastest). The User 20 may also choose the views desired from a pathway generator menu. For example, the User 20 may choose to display only the static and textual displays and those displays may be maximized upon activation of the system of the present invention.

Choosing an attribute may activate a read-only version of the concept editor. The event editor may also be activated by clicking on the event arrow corresponding to an event. Any text marked as a hyperlink (e.g., blue and underlined) will preferably bring up a Web browser to display information from the Internet when chosen by the user.

As illustrated in Fig. 13, Enter User Name and Password window 5100 is an example of a user login screen which may be used in the present invention. In this embodiment, a User 20 may be required to enter a user login ID and password. The user login ID and password may be maintained by a client. A User 20 may press the OK button 5102 to communicate the entered user login ID and password to the client. The

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user login ID and password may be authenticated against security information stored in a database of the client..

In an embodiment of the present invention, a preferred static display of pathways is as shown in window 5110 in Fig. 14. Window 5110 may show feedback loops. Feedback loops may be points in pathways in which the pathway regulates itself by feeding back to an original point in the pathway. A User 20 may print the display to a file or directly to a printer. A User 20 may select concepts or events and view all of the data associated with the selected concept or event.

In an embodiment of the present invention, a preferred dynamic display of pathways is as shown in window 5120 in Fig. 15.

In an embodiment of the present invention, a preferred textual display is as shown in window 5140 and window 5130 in Fig. 16 and Fig. 17, respectively. As shown in windows 5130 and 5140, the textual display may display the user entered initial conditions including user name, pathway type, context, stimulus, exclusions and endpoints. The textual display may be outputted to a file and/or printed. The textual display may be organized in several different ways. For example, window 5130 shows the textual display ordered by the pathway that occurs in the static display. In contrast, window 5140 shows the textual display ordered as they occur in time steps.

Fig. 18 illustrates a Pathway Generator window 5150 of the present invention. The Pathway Generator window 5150 may be the main window after the Enter User Name and Password window 5100. The user login ID entered in the Enter User Name and Password window 5100 may be displayed may be displayed in box 5151 of the Pathway Generator window 5150. In the Pathway Generator window 5150, a User 20 may choose from at least two pathway types. For example, a User 20 may select

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Forward Pathway radio button 5152 to execute a forward pathway. Alternatively, a User 20 may select Reverse Pathway radio button 5153 to execute a reverse pathway. A User 20 may select a cell from the Cell Type list box 5154 by highlighting a cell and selecting the Enable button 5155. This may load the cell into the Cell Type window 5156. In an embodiment of the present invention, a cell must be chosen and only one cell may be chosen.

A User 20 may choose a stimulus or stimuli when running the forward pathway generation and may choose a stimulus/stimuli when running the reverse pathway generation. A User 20 may choose a stimulus or stimuli by selecting the Stimuli Search button 5158. Upon selecting the Stimuli Search button 5158, a User 20 may be presented with the Search for Stimulus window 5180 shown in Fig. 19. If a User 20 selects a stimulus/stimuli, the corresponding information may be displayed in the Stimulus window 5157. A User 20 may choose to delete a given stimulus by selecting a stimulus in the Stimulus window 5157 and subsequently selecting the Delete button 5159.

A User 20 may choose a concept knockout when running the forward pathway generation or reverse pathway generation. A User 20 may choose to search all available concept knockouts by selecting the Knockout Search button 5161. Upon selecting the Knockout Search button 5161, a User 20 may be presented with the Search for Concepts window 5190 shown in Fig. 20. If a User 20 selects a knockout concept, the corresponding information may be displayed in the Concept Knockout window 5160. A User 20 may choose to delete a given concept knockout by selecting a concept knockout in the Concept Knockout window 5160 and subsequently selecting the Delete button 5162.

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A User 20 may choose a pathway endpoint when running the forward pathway generation or reverse pathway generation. A User 20 may choose to search all available pathway endpoints by selecting the Endpoint Search button 5164. Upon selecting the Knockout Search button 5164, a User 20 may be presented with the Search for Pathway Endpoints window 5200 shown in Fig. 21. If a User 20 selects a pathway endpoint, the corresponding information may be displayed in the Pathway Endpoints window 5163. A User 20 may choose to delete a given pathway endpoint by selecting a pathway endpoint in the Pathway Endpoints window 5163 and subsequently selecting the Delete button 5165.

A User 20 may choose to exclude certain information from the pathway generation by selecting the Event Exclusion Criteria button 5166. If a User 20 selects the Event Exclusion Criteria button 5166, an Event Exclusion Criteria window 5210 may be presented as shown in Fig. 22. Any criteria entered by a User 20 in the Event Exclusion Criteria window 5210 may be displayed in the Event Exclusion Criteria box 5167. A User 20 may clear the Event Exclusion Criteria box 5167 by selecting the Clear button 5168. A User 20 may execute the pathway based on the selected information by selecting the Run CellTek button 5169. A User 20 may clear all information in the Pathway Generator window 5150 by selecting the Clear All button 5170. A User 20 may exit the Pathway Generator window 5150 by selecting the Exit CellTek button 5171. In a preferred embodiment, a User 20 may save the pathway conditions from a given pathway to a file. A User 20 may also open information previously saved in the Pathway Generator window 5150.

As shown in Fig. 19, a User 20 may enter a full text search for information by entering the full text into the Query String field 5181, selecting the Full String Search

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check box 5182, and subsequently selecting the Search button 5183. A User 20 may also enter partial string searches by deselecting the Full String Search check box 5182, entering the partial search information into the Query String field 5181, and subsequently selecting the Search button 5183. Upon selection of the Search button 5183, the results of the search are displayed in the Result box 5184. A User 20 may choose to use the searched information by selecting the given information in the Result box 5184 and subsequently selecting the OK button 5185. The selected information will then be moved to a corresponding location in the Pathway Generator window 5150. A User 20 may also select the Cancel button 5186 to return to the Pathway Generator window 5150. When a User 20 is finished using the Search for Stimulus window 5180, a User 20 may select the Exit button 5187 to return to the Pathway Generator window 5150.

Fig. 22 illustrates an Event Exclusion Criteria window 5210. A User 20 may expand the Event Exclusion Criteria window 5210 by selecting the More button 5211. Expansion of the Event Exclusion Criteria window 5210 allows the User 20 to enter more information. A User 20 may select a pre-defined field to exclude data by selecting from a list in the Field list 5212. The pre-defined fields may include references, experiment temperature, assays and species. A User 20 may also choose a selection in the Qualifier list 5213. The Qualifier list 5213 may include equal to, not equal to, less than, greater than, less than or equal to, or greater than or equal to. The options presented in the Values box 5214 may be dependent upon the selection made in the Field list 5212. A User 20 may search for values in the Database 80 using the Search button 5215. The Operator button 5216 may be a link between multiple rows of event exclusion criteria and may include AND or OR. A User 20 may enter information by selecting the OK button 5217, the Event Exclusion Criteria

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window 5210 may close and entered information may be transferred to the Pathway Generator window 5150. Alternatively, a User 20 may close the Event Exclusion Criteria window 5210 without transferring the information by selecting the Cancel button 5218.

Fig. 23 illustrates a Concept Editor window 5220. The Concept Editor window 5220 may be displayed when a User 20 edits concept information or view information from a static display as a read-only display. A User 20 may search for a reference by selecting a Search button 5221. The Search button 5221 may open a Search for Reference window 5260. Upon selection of a reference, the reference information may be displayed in an Active Reference box 5222. If a reference is active (e.g., the reference is relevant to all entered information), the reference may be indicated with a check mark in the Active check box 5223. In an embodiment of the present invention, User 20 must have an active reference to enter information into the Database 80. A User 20 may search for concepts by selecting the Search button 5270. The Search button 5270 may present the Search for Concepts window 5190 (Fig. 20). Alternatively, a User 20 may enter a new concept by selecting a New button 5225. Upon selection of the New button 5225, the New Concept Name window 5280 may be presented. A User 20 may delete a concept by selecting a Delete button 5226. A User 20 may indicate if the concept is transportable by enabling the Transportable check box 5256. A User 20 may select a class for a given concept by entering the class from a list of classes presented in Class box 5227.

A User 20 may accept the selected class by selecting the Accept button 5228. A User 20 may also view a detailed listing of all class information by selecting a Class button 5229. Upon selection of the Class button 5229, a Class Concepts Hierarchies window 5290 may be presented. As shown in Fig. 27, the Class Concepts Hierarchies

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window 5290 may show all of the class information. A User 20 may refresh the Concept Editor window 5220 by selecting the Refresh button 5230. A User 20 may also Add, Edit, or Delete names by selecting button 5231, 5232, or 5233, respectively, subsequent to selecting a given name in the Names window 5235. Upon selection of a name in the Names window 5235, the selected name may be displayed in the Name Text box 5234 and a drop down list may be displayed with the name type below the Name Text box 5234.

A User 20 may cancel the active name by selecting the Cancel button 5255. A User 20 may choose to associate a concept with an organism by selecting the Add/Change button 5236, if an active relationship exists. Corresponding organism information may be displayed in the Organism window 5238. A User 20 may delete information by selecting the information in the Organism window 5238 and subsequently selecting the Delete button 5237. A User 20 may expand the concept from another concept by selecting the Add/Change button 5239, if an active relationship exists. The expanded information may be displayed in the Expands Concept window 5241. A User 20 may delete information by selecting given information in the Expands Concept window 5241 and subsequently selecting the Delete button 5241. A User 20 may join a concept with another concept by selecting the Add/Change button 5242, if an active relationship exists. The joined information may be displayed in the Joins Concept window 5244. A User 20 may delete information by selecting the given information in the Joins Concept window 5244 and subsequently selecting the Delete button 5243.

A User 20 may add primitive attributes to a concept by selecting the Primitive button 5245. Upon selection of the Primitive button 5245, a Primitive Attributes window 5300 may be presented, as shown in Fig. 28. A User 20 may choose to search

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references by selecting the References button 5246. Upon selection of the References button 5246, a References window 5360 may be presented, as shown in Fig. 34. A User 20 may choose to add contains information to a concept (e.g., a container) by selecting the Contains button 5247. Upon selection of the Contains button 5247, the Contains window 5310 may be presented, as shown in Fig. 29. A User 20 may choose to add database information to a concept by selecting the DB/UI button 5248. Upon selection of the DB/UI button 5248, the External Databases window 5370 may be presented, as shown in Fig. 35. A User 20 may choose to add anatomic attributes to a concept by selecting the Anatomic button 5249. Upon selection of the Anatomic button 5249, an Anatomic Attributes window 5330 may be presented, as shown in Fig. 31. A User 20 may choose to add scope notes to a concept by selecting the Scope button 5250. Upon selection of the Scope button 5250, a Scope Notes window 5380 may be presented, as shown in Fig. 36. A User 20 may choose to add molecular attributes to a concept by selecting the Molecular button 5251. Upon selection of the Molecular button 5251, a Molecular Attributes window 5340 may be presented, as shown in Fig. 32. A User 20, may choose to add editorial notes to a concept by selecting the Editorial button 5252. Upon selection of the Editorial button 5252, an Editorial Comments window 5390 may be presented, as shown in Fig. 37. A User 20 may choose to add reagents to a concept by selecting the Reagents button 5253. Upon selection of the Reagents button 5253, a Reagents window 5350 may be presented, as shown in Fig. 33. A User 20 may choose to associate the active concept with an event by selecting the Events button 5254. Upon selection of the Events button 5254, an Events Editor window 5400 may be presented, as shown in Fig. 38.

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Fig. 24 illustrates a Search for Reference window 5260. The Search for Reference window 5260 may allow a User 20 to perform partial searches on reference information. A User 20 may enter author information in the Author text box 5262. A User 20 may enter reference titles in the Title text box 5263. A User 20 may enter publication year in the Year text box 5264. A User 20 may enter the distinct reference PMID in the PMID text box 5265. A User 20 may select the Search button 5266 to search based on the entered information. All information queried from the Database 80 corresponding to the given criteria may be displayed in the Results window 5268. A User 20 may select a reference in the Results window 5268. Upon selection of a reference in the Results window 5268, the full reference information may appear in the Full Reference Information window 5269. A User 20 may choose to make a reference active by selecting the Make Active button 5267.

Fig. 25 illustrates a Search for Concept Name window **5270**. The Search for Concept Name window **5270** may operate in a manner similar to the Search for Stimulus window **5180** (Fig. 19), the Search for Concepts window **5190** (Fig. 20), or the Search for Pathway Endpoints window **5200** (Fig. 21).

Fig. 26 illustrates a New Concept Name window 5280. The New Concept Name window 5280 may be used to enter a new concept. An active concept may be displayed in the Concept window 5281. A User 20 may enter a new concept name in the Concept box 5282. A User 20 may search for duplicate names by pressing the Validate button 5283. If a name is not a duplicate, the Validate button 5283 may not be selected by a User 20. If a name is a duplicate, an error message may be displayed to the User 20. A User 20 may accept the new name by selecting the Accept button 5284. A User 20 may exit the New Concept Name window 5280 by selecting the Exit button 5285.

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Fig. 27 illustrates a Class Concepts Hierarchies window 5290. The Class Concepts Hierarchies window 5290 may be used to establish a class relationship to a concept. A User 20 may view the various class by browsing through the hierarchies in the Hierarchies window 5294. A User 20 may choose a relation by selecting the Relations button 5291. A User 20 may make the relationship active by selecting the Active button 5292. A User 20 may refresh the Hierarchies window 5294 by selecting the Refresh button 5293.

Fig. 28 illustrates a Primitive Attributes window 5300. The Primitive Attributes window 5300 may be used to assign primitive attributes to a concept. For example, a User 20 may assign values for display to a concept. A User 20 may also indicate whether a concept is a stimulus. A User 20 may add the entered primitive attributes to the concept by selecting the Add button 5301. A User 20 may refresh the Primitive Display box 5305 by selecting the Refresh button 5303.

Fig. 29 illustrates a Contains window 5310. The Contains windows 5310 may be used for showing containment of concepts in cellular context (e.g., cells, cell structures, etc.). A User 20 may search for concepts to add to a cellular context. A User 20 may initiate a search by selecting the Search button 5311. Upon selecting the Search button 5311, a User 20 may be presented with a search dialog and/or search results. Concept search results may be selected by a User 20 for containment in the cellular context. Such selected concepts may be displayed in the Contains Concept box 5316. Similarly, a User 20 may select concepts to exclude from the cellular context. Such concepts may be displayed in the Excludes Concepts box 5319. A User 20 may also locate concepts to contain in or exclude from the cellular contexts by utilizing a batch query as discussed in reference to Fig. 30.

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Fig. 30 illustrates a Query window 5320. A User 20 may utilize the Query window 5320 to execute a batch import of concepts in the Contains Concept box 5316 or the Excludes Concepts box 5319. A User 20 may choose to include or exclude an organism by selecting Include button 5321 or Exclude button 5322, respectively. A User 20 may choose to include or exclude an anatomic by selecting Include button 5323 or Exclude button 5324, respectively. After selecting the appropriate include or exclude radio button, a User 20 may choose organism specific information by selecting the Ouery button 5325, selecting the desired concepts, and selecting the Add button 5326. Similarly, a User 20 may choose anatomic information by setting up a search by selecting the desired contextual limitations (e.g., developmental stage, organ, tissue, cell type, etc.) and search term connectors (e.g., AND, OR, etc.). A User 20 may execute the search by selecting the Query button 5327. A User 20 may select class information by selecting the Query button 5328 to find concepts, selecting the desired concepts and then selecting the Add button 5329. If a User 20 wishes a class to be displayed with all sub-classes or inherited classes, a User 20 may select the Expand check box 5321a. A User 20 may execute individual queries for an organism, anatomic or class by selecting the corresponding query button. A User 20 user may execute a combination query for an organism, anatomic and/or class by selecting the Combination Query button 5321b. The Display box 5321c may display all information resulting from any searches or queries. A User 20 may send information displayed in the Display box 5321c to the Contains Concept box 5316 or the Excludes Concepts box 5319 by selecting the information and selecting the To Contains button 5321e. A User 20 may search for general concepts by selecting the Search button 5321d.

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Fig. 31 illustrates an Anatomic Attributes window 5330. A User 20 may use the Anatomic Attributes window 5330 to associate anatomic information (e.g., organ, tissue, cell line, etc.) with a concept. The Top Display window 5331 displays information regarding an active reference. An active reference is a reference associated with a given concept. When new information is added to a concept, a new reference may be added. New reference information may be displayed in an Active ID Reference box 5333. Line item information for a new reference may be displayed in the Line Item Display box 5332. The Add/Change buttons 5334, 5336, 5338 and 5335 may be used to add and/or change information regarding developmental stage, organ, tissue and/or cell type. When a User 20 selects the Accept button 5339, information entered through the Anatomic Attributes window 5330 is validated and entered into the Database 80.

Fig. 32 illustrates a Molecular Attributes window 5340. A User 20 may use the Molecular Attributes window 5340 to associate molecular information with a concept. The Top Display window 5341 may display information regarding an active reference. The Member of Gene/Protein Family window 5342 displays information regarding membership of a reference or concept in a gene or protein family. The Has a Prototype Homolog window 5343 displays information regarding prototype homologs a reference or concept may have. A User 20 may select Domains radio button 5344, Motifs radio button 5345, Post-Translational Modifications radio button 5346, Activated By radio button 5347 or Inhibited By radio button 5348 to display corresponding Active ID information in Active ID Display box 5349. The References box 5349e may display more detailed reference information regarding the Active ID information corresponding to select Domains radio button 5344, Motifs radio button 5345, Post-Translational

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Modifications radio button **5346**, Activated By radio button **5347** or Inhibited By radio button **5348**.

Fig. 33 illustrates a Reagents window 5350. Top Display window 5351 may display information regarding an active reference. When a User 20 selects the Add button 5355, the User 20 may be prompted to enter a reagent name. Upon entering a reagent name, the reagent name may be displayed in Middle Display window 5352. A User 20 may add references associated with each reagent in Bottom Display window 5353 by selecting a given reference displayed in Bottom Display window 5353 and subsequently selecting the Add button 5355.

Fig. 34 illustrates a References window **5360**. The References window **5360** may display references associated with a given concept. A User **20** may choose to activate a reference by selecting a reference from a list of references and subsequently selecting an Add Active button **5364**. In the embodiment illustrated in Fig. 34, a User **20** may select a To 'Search' button **5363** to jump to the Search for Reference window **260** (Fig. 24).

Fig. 35 illustrates an External Databases window 5370. A User 20 may utilize the External Databases window 5370 to identify external databases for use by the Inference Engine 14. These external databases may be utilized by the Inference Engine 14 in conjunction with the Database 80. A User 20 may also utilize the External Databases window 5370 to identify unique identifiers associated with a concept.

Fig. 36 illustrates a Scope Notes window 5380. A User 20 may associate scope notes with a concept by entering the scope notes and selecting the Add button 5386. When a User 20 enters scope notes, selects the Add button 5386 and selects the Exit button 5382, the entered scope notes may be displayed in the Upper Display window 5387. Line item information may be displayed in Lower Display window 5381. A User

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20 may select a line item in Lower Display window 5381. Subsequently, a User 20 may select an Edit button 5385 to edit the selected line item. A User 20 may also select a Delete button 5384 to delete the selected line item.

Fig. 37 illustrates an Editorial Comments window 5390. A User 20 may associate editorial comments with a concept by entering the editorial comments and selecting the Add button 5396. When a User 20 enters editorial comments, selects the Add button 5396 and selects the Exit button 5392, the entered editorial comments may be displayed in the Upper Display window 5397. Line item information may be displayed in Lower Display window 5391. A User 20 may select a line item in Lower Display window 5391. Subsequently, a User 20 may select an Edit button 5395 to edit the selected line item. A User 20 may also select a Delete button 5394 to delete the selected line item.

Fig. 38 illustrates an Event Editor window 5400. A User 20 may utilize the Event Editor window 5400 to enter information associated with events. Active Reference window 5401 may display an active reference. Event window 5402 may display the list of events associated with the active reference. A User 20 may add an event by selecting the New Event button 5400a. A User 20 may search for an event by selecting Search Event button 5408. Upon selection of the Search Event button 5408, the Search for Event Name window 5405 may be presented. Names window 5403 may display names and types of events. A User 20 may add or edit the names in Names window 5403 by entering information into Text window 5404. In an embodiment of the present invention, a User 20 must enter relevant information into the Requires window 5405 and the Produces window 5406. A User 20 may associate an event with a controversy flag by selecting the Controversy Flag check box 5407. A User 20 may select concepts by which the event is inhibited by selecting the Add button 5400b. A

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User 20 may also select where an event occurs by adding location information to a Cellular Location window 5400c. A User 20 may select a variety of different attributes to be associated with the event by selecting Constants button 5400d, Attributes button 5400e, References button 5400f, Editorial button 5400g, Experimental button 5400h, Containers button 5400i, DB/UI button 5400j, or Scope button 5400k.

Upon selecting Constants button 5400d, a User 20 may be presented with Biochemical Constants window 5410 (Fig. 40). Upon selecting Attributes button 5400e, a User 20 may be presented with Event Attributes window 5420 (Fig. 41). Upon selecting Experimental button 5400h, a User 20 may be presented with Experimental Conditions window 5430 (Fig. 42). Upon selecting References button 5400f, a User 20 may be presented with References window 5360 (Fig. 34). Upon selecting Editorial button 5400g, a User 20 may be presented with Editorial Comments window 5390 (Fig. 37). Upon selecting DB/UI button 5400j, a User 20 may be presented with External Databases window 5370 (Fig. 35). Upon selecting Scope button 5400k, a User 20 may be presented with Scope Notes window 5380 (Fig. 36). Upon selecting Search Concepts button 5400l, a User 20 may be presented with Search for Concept Name window 270 (Fig. 25).

Fig. 39 illustrates a Search for Event Name window **5405**. The Search for Concept Name window **5405** may operate and be utilized in a manner similar to the Search for Stimulus window **5180** (Fig. 19), the Search for Concepts window **5190** (Fig. 20), or the Search for Pathway Endpoints window **5200** (Fig. 21).

Fig. 40 illustrates a Biochemical Constants window **5410**. Active Reference window **5411** may display an active reference. A User **20** may enter a maximum velocity in Vmax window **5412**. A User **20** may enter a Michaelis constant in Km window **5413**.

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A User 20 may enter an equilibrium constant in Keq window 5414. A User 20 may enter a dissociation constant in Kd window 5415. A User 20 may indicate that all constants are known by selecting Completed Constants check box 5416. Kinetic Display check box 5417 may be selected to indicate that the forward and reverse kinetic constants are known. A User 20 may enter a reverse kinetic constant in Reverse Rate window 5418. A User 20 may enter a forward kinetic constant in Forward Rate window 5419. Upon selection of Accept button 5410a, information entered in Biochemical Constants window 5410 may be validated and entered in Database 80.

Fig. 41 illustrates an Event Attribute window 5420. An event name may be displayed in Event Name window 5421. A User 20 may add attributes to the event in the Has Attributes window 5422. A User 20 may add test attribute conditions to the event by selecting Add button 5420a. A User 20 may add information that modifies an attribute of an event by selecting the Add button 5420b. A User 20 may add information that applies to an event by selecting the Add button 5420c associated with the Applies Process window 5420d.

Fig. 42 illustrates an Experimental Conditions window 5430. Active Reference window 5431 may display an active reference. Line items for all of the experimental condition information and associated reference IDs may be displayed in Experimental Conditions Display window 5432. When a User 20 selects a line item in Experimental Conditions Display window 5432, appropriate reference information may be displayed in Reference window 5433. When a User 20 searches for an assay using Add/Change button 5430b, a corresponding assay name may be displayed in Assay Name window 5434. When a User 20 searches for a preparation type using Add/Change button 5430c, a corresponding event preparation type may be displayed in Sample Preparation Type

window 5435. Assay Description window 5436 may display an assay description. Assay Buffer window 5437 may display assay buffer information. Temperature window 5438 may display the experimental temperature. The information entered into the Experimental Conditions window 5430 may be verified and entered into the Database 80 by selecting the Accept button 5430a.

Fig. 43 illustrates an Excluded window 5440. The Excluded window 5440 may be displayed upon selection of the Containers button 5400i (Fig. 38). A User 20 may utilize the Excluded window 5440 to select containers from which the event is excluded. A User 20 may search for a given container upon selecting Search button 5440a. A User 20 may add a given container by selecting the Add button 5440b.

STL Shell

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Examples are now presented for demonstration of grammar and representation only. Statements may have no basis in scientific fact.

Attributes: Attributes are used to annotate concepts and events. They are placeholders that represent information that can be assigned to a concept or an event. The mass of an object, the color of a protein on the screen, and all of the parameters that define how an object will move are all described by attributes.

The formal specification for defining an attribute is as follows:

ATTRIBUTE <attribute name > IS [ANY] [INTEGER | REAL | TEXT | OF] [<value list >] [FROM [EXACTLY] <number > TO [EXACTLY] <number >] [INCLUSIVE].

The grammar in square brackets [] is optional.

Some specific examples of definitions (with explanations):

Attribute "mass" is any real.

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This statement defines an attribute called "mass" that can equal any real (decimal) number.

Attribute "abstract" is any text.

This statement defines an attribute called "abstract" that can equal any text.

Attribute "location" is any of "membrane-bound", "extracellular", "cytoplasmic".

This statement defines an attribute called "location" that can have any one of the values "membrane-bound", "extracellular" or "cytoplasmic".

Attribute "color" is any integer from 1 to 10 inclusive.

This statement defines an attribute called "color" that can have any integral value greater then or equal to 1, and less than or equal to 10. The "inclusive" keyword means that both the from-value and the to-value can be equal to their limits- a shorthand way of stating from exactly 1 to exactly 10.

Attribute "weight" is any real from 0.3.

This statement defines an attribute called "weight" that can have any real value greater than 0.3.

Attribute "height" is any real to exactly 14.3.

This statement defines an attribute called "height" that can have any real value less than or equal to 14.3.

Some of the attributes will be used by the Simulation Module 10 to determine whether or not a reaction will proceed. Most attributes will be used by the dynamic graphic display of the Simulation Module 10 to determine how the events will be represented on the screen. Attributes can also be used to store information about the concepts for hyperlinks and other data displays.

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The following is a list of attributes that may be directly used by the system and method of the present invention.

Attribute Name, Purpose

- <u>Substance</u>: Any concept that has this attribute will be listed as a substance in certain lists in the Concept, Event and Pathway wizards. Does not require any particular value to be assigned.
- <u>Process</u>: Any concept that has this attribute will be listed as a process in the Process list in the Event wizard. Does not require any particular value to be assigned.
- <u>Structure</u>: Any concept that has this attribute will be listed as a "structure" (cell or cellular component) in certain lists in the Concept, Event and Pathway wizards. Does not require any particular value to be assigned.
 - <u>Duration</u>: Assigned to an event, determines the length of time in seconds the event requires to proceed. May be a decimal number.
 - Graphicshape: Assigned to a concept, determines the shape of the concept in graphical presentation. Enumerated list with the following elements: none, triangle, square, circle.
 - <u>Graphicsize</u>: Assigned to a concept, determines the relative size of the concept in graphical presentation. Number from 1 to 100.
 - <u>Graphiccolor</u>: Assigned to a concept, determines the color of the concept in graphical presentation. Long integer in RGB format.
 - Location: Assigned to a concept, determines the starting cellular location of the concept in the dynamic graphical presentation. Enumerated list with the following elements: MB (membrane bound), Cyt (cytoplasmic), Comp (complexed), Nuc (nuclear), EC (extracellular) and ER (endoplasmic recticulum).
- <u>Mobile</u>: Assigned to an event, determines which reactant(s) in the event will move in the dynamic graphical presentation. Consists of a string of 'Y' or 'N' characters, 'Y' means will move and 'N' means won't move. The characters are given in the order that the reactants are listed.
- <u>Postevent</u>: Assigned to an event, determines which reactant(s) in the event will remain visible after the event is over in the dynamic graphical presentation. Consists of a string of 'Y' or 'N' characters, 'Y' means will be visible and 'N' means won't be visible. The characters are given in the order that the reactants are listed.
 - <u>Stimulus</u>: Any concept that has this attribute will be listed as a stimulus in the Stimulus list in the Pathway wizard. Does not require any particular value to be assigned.

Every substance or process is represented by a concept. Concepts can be defined as special types of other concepts, and can contain other concepts. Concepts can be assigned attributes, and may assign values to their attributes.

40 <u>Concepts:</u> The formal specification for defining a concept is as follows:

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CONCEPT <concept name> [EXPANDS <base concept>[;]] [CONTAINS <concept list>[;]] [EXCLUDES <concept list>[;]] [JOINS <concept list>[;]] [HAS <attribute list>[;]] [SETS <attribute assignment list>].

The grammar in square brackets [] is optional.

Some specific examples of definitions (with explanations):

Concept "object" has "mass", "size".

This statement defines a concept called "object" to which the attributes "mass" and "size" have been assigned.

Concept "protein" expands "object".

This statement defines a concept called "protein" as a specific type of "object". The "protein" concept inherits the attributes of "object", therefore it has attributes "mass" and "size" associated with it automatically.

Concept "TNFR" expands "protein"; has "abstract"; sets "mass"=30, "abstract" = "abstract of TNFR".

This statement defines a concept called "TNFR" as a specific type of "protein". It inherits the attributes of a "protein", and therefore automatically has "mass" and "size" attributes. It gives itself an "abstract" attribute (documentation of the protein). It sets its "mass" attribute to 30 (for the time being, we will use normalized values for physical attributes), and its "abstract" to the text "abstract of TNFR".

Concept "generic cell" expands "object"; contains "nucleus", "mitochondria", "cell membrane"; excludes "RAF"; sets "size"=40.

This statement defines a concept called "generic cell" as a specific type of "object". It inherits "mass" and "size" attributes from "object". It defines itself as a wrapper around the concepts "nucleus", "mitochondria", and "cell membrane", and sets its "size" to 40. It prevents the inheritance of the "RAF" concept from any of its child

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concepts. Concepts that contain other concepts do not inherit attributes from the concepts they contain.

Concept "Active FADD" expands "protein"; joins "active", FADD".

This statement defines a concept called "Active FADD" as a specific type of protein. It inherits the attributes of a protein, and joins together the concepts "active" and "FADD". Concepts that represent conjoined concepts inherit attributes from the concepts they contain. Conjoined concepts currently have little utility in the present invention; activated proteins are represented as inheriting from the inactivated forms:

Concept "active FADD" expands "FADD".

The events determine which concepts react, what concepts are produced as a result, the processes that occur within the reaction, and what determines whether or not the reaction will proceed.

15 <u>Events:</u> The formal specification for an event is as follows:

EVENT <event name> REQUIRES <concept list>[;] [TESTS <attribute> OF <concept> {EQUAL TO | LESS THAN | GREATER THAN} <attribute> OF <concept>[;]] [APPLIES <concept> FROM <concept> TO <concept>[;]] [PRODUCES <concept list>[;]] [INHIBITED BY <concept list>[;]] [PRESENT IN <concept list>[;]] [ABSENT IN <concept list>[;]] [HAS <attribute list>[;]] [SETS <attribute assignment list>].

The grammar in square brackets [] is optional.

25 Some examples of definitions (with explanations):

Event "Three molecules of TNF binds to and trimerizes TNFR" Requires "TNF", "TNF", "TNFR", "TNFR", "TNFR", "TNFR"; Applies "binds to" from "TNF" to "TNFR"; Applies "trimerizes" from "TNF" to "TNFR"; Produces "Trimer of TNFR"; Has "duration"; Sets "duration"=10.

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This statement defines an event that requires three instances of the "TNFR" concept and three instances of the "TNF" concept to proceed. It produces the concept "Trimer of TNFR", and requires a "duration" of 10 to do so. It performs the reaction by applying the "binds to" and "trimerizes" concepts from "TNF" to "TNFR". (The purpose of the 'applies' clause is to break the reaction into pieces to describe the process to the static presentation engine.)

Event "GTP displaces GDP and activates heterotrimeric Gs protein" Requires "GTP", "GDP-bound heterotrimeric Gs protein"; Tests "concentration" of "GTP" greater than "concentration" of "GDP"; Produces "active heterotrimeric Gs protein", "GDP".

This statement defines an event that requires "GTP" and "GDP-bound heterotrimeric Gs protein" to proceed. It produces the concepts "GDP" and "active heterotrimeric Gs protein". It tests that the attribute "concentration" of concept "GTP" is numerically greater than the attribute "concentration" of concept "GDP", and if so, permits the reaction to take place. No processes are specified, so if this event occurs it will not be fully animated. No duration is specified, so the animation engine will assign it a default reaction time of 5 seconds.

Event "Adenylyl cyclase converts ATP to cAMP and Pi" Requires "Adenylyl cyclase", "ATP"; Applies "converts ATP to" from "adenylyl cyclase" to "ATP"; Present In "generic cell"; Produces "cAMP", "Pi".

This statement defines an event that requires "adenylyl cyclase" and "ATP" to proceed. It produces the concepts "cAMP" and "Pi". It will only occur in a "generic cell" context. It performs the reaction by applying the "converts ATP to" concept from "adenylyl cyclase" to "ATP". (This may be an area of some contention as to which

concept "performs" the action and which concept it is "performed on". Naturally, the most scientifically appropriate definition of the process should be used if it is known.)

Event "TRADD binds to and activates TRAF2"
Requires "active TRADD", "TRAF2";
Inhibited by "FADD";
Absent In "liver cell";
Applies "binds to" from "TRADD" to "TRAF2";
Applies "activates" from "TRADD" to "TRAF2";
Produces "active TRAF2".

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This statement defines an event that requires "active TRADD" and "TRAF2" to proceed. It produces that concept "active TRAF2". It performs the reaction by applying "binds to" and "activates" from "TRADD" to "TRAF2". The reaction will not occur if the concept "FADD" is present. The reaction will not occur in a liver cell (but will occur in every other type of cell).

The following tables may be used by the system and method of the present invention. They are preferably stored in structure form in memory. They are preferably read from and written to a structured storage document.

Data Tables:

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The ANYVALUE structure is a VARIANT-like type that provides a transparent means of storing information of different types.

Name	Type	Description
Lnum	long	Part of an anonymous union, stores a long integer value.
Fnum	double	Part of an anonymous union, stores a double floating point value.
Index	long	Part of an anonymous union, stores an index into a string table or enumerated value table.
Error	long	Part of an anonymous union, stores an error code.
Hasvalue	bool	Indicates whether the union contains a valid value.
Туре	char	Indicates the type of the data contained within the union.

The ANATOMIC table holds the anatomical information.

Name	Type	Description
Anatomic_Id	long	Primary key for this table. Anatomic identifier.
Tissue	long	Identifies the tissue associated with the concept.
Reference_Id	long	Identifies the reference.
CellType	long	Identifies the cell type.
MoleculesPerCell	double	Identifies the molecules per cell.
CellLine	text	Identifies the cell line
DevelopmentalStage	long	Identifies the developmental stage.
Organ	long	Identifies the organ
AnatExpression_Id	long	Identifies the anatomical expression.

The ANATOMICEXPRESSION table holds the types of anatomical expression information.

Name	Type	Description
AnatExpression_ID	long	Primary key for this table. Anatomic identifier.
Tissue	text	Identifies the anatomical expression type (e.g. Protein, RNA, etc.)

The APPLIES table holds the applies process information.

Name	Type	Description
Apply_Id	long	Primary key for this table. Identifier for the table.
Apply_Concept	long	Identifies the concept that is applied.
From_Concept	long	Identifies the concept that the relationship comes from.
To_Concept	long	Identifies the concept the relationship points to.

The ASSAYNAME table holds the names of the assays used in the generation of the events.

Name	Type	Description
Assay_Id	long	Primary key for this table. Applies identifier.
Name	Text	Identifies the name of the assay.

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The ATTRIBUTEENUMVALUE table holds all possible enumerated values for all enumerated list attributes. The offset within this table identifies which element of the enumerated list an enumerated list attribute represents.

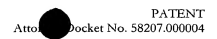
Name	Type	Description
AttributeENum_Id	long	Composite key for this table with Name.
		Attribute type identifier.
Name	text	Composite key for this table with
14.1		AttributeENum_Id. Name of the attribute (e.g.
		shape, cellular location, etc.)

The ATTRIBUTES table holds the attribute definitions.

Name	Type	Description
Attribute_Id	long	Primary key for this table. Attribute identifier.
Name	text	Name of the attribute.
Туре	long	Identifies the type of the attribute. (e.g. integer, real, enumerated list, etc.)
Bounds	char	Identifies the boundary limits of the attribute, if it is a number.
Upperbound	long	Stores the upper bound of the attribute if it is numerical.
Lowerbound	long	Stores the lower bound of the attribute if it is numerical.
Upperexact	long	Indicates whether the upper bound is inclusive if the attribute is numerical.
Lowerexact	long	Indicates whether the lower bound is inclusive if the attribute is numerical.

The AUTHORLINK table holds the links between authors and references.

Name	Type	Description
AuthorLink_Id	long	Primary key for this table. Author link identifier.
Author_Id	long	Identifies the author.
Reference_Id	long	Identifies the reference.



The AUTHORS table holds the author information.

Name	Type	Description
Author_Id	long	Primary key for this table. Author identifier.
Name	Text	Identifies the name of the author.

The CLASSES table holds the hierarchical relationships between concepts.

Name	Type	Description
Concept_Id	long	Primary key for this table. Concept identifier.
Parent	long	Parent of the concept.
ClassType_Id	long	Class type identifier.

5 The CLASSTYPE table holds the hierarchical relationship types.

Name	Type	Description
ClassType_Id	long	Primary key for this table. Class type identifier.
Description	long	Describes the class type.

The COMMENT table holds the comment information provided by the editors.

Name	Type	Description
Comment_Id	long	Primary key for this table. Comment identifier.
CommentType_Id	long	Identifies the type of comment. (e.g. editor comments, scope notes, user notes, etc)
CommentRefType_Id	long	Identifies the comment reference type as "concept" or "event".
Id	long	Identifies the concept or event Id that is associated with the given comment.
Comment	text	Editor comments.

The COMMENTREFTYPE table holds the comment reference type

10 information.

Name	Type	Description
CommentRefType_Id	long	Primary key for this table. Comment reference type identifier.
Name	text	Name of the comment reference type. (Concept or Event)

The COMMENTTYPE table holds the comment type information.

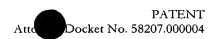
Name	Type	Description
CommentType_Id	long	Primary key for this table. Comment reference type identifier. (e.g. editor comments, scope notes, user notes, etc)
Name	text	Name of the comment type.

The COMPAREENUM table holds the enumerated list values for attribute comparison.

Name	Туре	Description
Compare_Id	Long	Primary key for this table. Comparison identifier. (e.g. equal to, less than, not greater than, etc)
Name	Text	Name of the comparison type.

The CONCEPTATTRIBUTE table holds the concept attribute information.

Name	Туре	Description
Concept_Id	long	Composite key for this table with Attribute_Id. Concept identifier.
Attribute_Id	long	Composite key for this table with Concept_Id. Attribute identifier.
StoredValue	text	Identifies the value of the concept's attribute.
HasValue	bool	Concept attribute indicator.
Туре	long	Identifies the attribute type.



The CONCEPTS table holds the concept definitions.

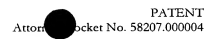
Name	Type	Description
Concept_Id	long	Composite key for this table. Concept identifier.
User_Id	long	Composite key for this table. User identifier.
Expands	long	Indicates another concept that the concept is based on.
Version_Id	long	Indicates the version of the concept for documentation purposes.
Organism	long	Indicates the organism the concept was present in.
Transportable	bool	Indicates whether the concept has the transportable feature.
Homolog_Of	long	Indicates what homolog(s) the concept may have.
GeneProteinFamily	long	Indicates concept membership in a gene or protein family.
Homolog_Flag	long	Indicates whether the concept is a homolog of another concept.

The DBUI table holds the reference information for outside databases and unique identifiers.

Name	Type	Description
DBUI_Id	long	Primary key for this table. DBUI identifier.
DBUIType	long .	Identifies the type of DBUI.
DB	long	Identifies the database type
UI	text	Identifies the UI type.

The DBUITYPE table holds the enumerated type values for the DBUI.

Name	Туре	Description
DBUIType_Id	Long	Primary key for this table. DBUI type identifier.
Name	Text	Name of the DBUI type. (e.g. string, numeric)



The EVENTATTRIBUTE table holds the relationships between the events and the event attributes.

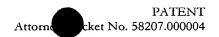
Name	Type	Description
Event_Id	long	Composite key for this table with Apply_Id. Event identifier.
Attribute_Id	long	Composite key for this table with Event_Id. Attribute identifier.
StoredValue	text	Actual value of the event attribute.
HasValue	bool	Event attribute indicator.
Туре	long	Identifies attribute type.

The EVENTS table holds the event definitions.

Name	Type	Description
Event_Id	long	Composite key for this table. Event identifier.
User_Id	Long	Composite key for this table. User identifier.
Controversy	long	Name or description of the event.
CellularLocation	long	Indicates whether this event has one or more Applies clauses (that define the processes in the event).
Version_Id	long	Indicates whether this event has one or more Tests clauses.
TransportLocation	long	Indicates whether this event has one or more attributes.
EquilibriumConstantEQ	long	Equilibrium constant.
DissociationConstantKD	long	Dissociation constant.
ForwardKineticRateConstantK1	long	Forward kinetic rate constant.
ReverseKineticRateConstantK1	long	Reverse kinetic rate constant.
MichaelKM	long	Michaelis-Menten constant
EnzymaticVMax	long	Enzymatic maximum velocity.
KineticDisplay	bool	Kinetic display indicator.
CompletedConstants	bool	Completed constant indicator.

The EVENTSPECIES table holds the event identifiers and the reference identifiers.

Name	Type	Description
Event_Id	long	Primary key for this table. Event identifier.
Organism	long	Organism identifier.



The EXPCONDITIONS table holds the experimental conditions in which the event was described.

Name	Type	Description
ExpCondition_Id	long	Primary key for this table. Experimental
		condition identifier.
Reference_Id	long	Identifier for references.
Assay	long	Identifier for assay.
Preparation_Type	long	Identifier for preparation type.
AssayProcess	text	Assay process description.
AssayBuffer	text	Assay buffer description.
TemperatureC	long	Temperature in degrees celcius.

The EXTERNALDB table holds the DB identifier and the associated database

5 name.

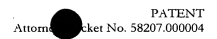
Name	Туре	Description
DB_Id	long	Primary key for this table. Database identifier.
Name	long	Name of the database.

The GENCONCEPT table holds the generic concept details.

Name	Type	Description
GenConcept_Id	long	Primary key for this table. Generic concept identifier.
GenConcept	long	Identifier for generic concepts.
Concept_Id	long	Identifier for concepts.
Type_Id	long	Identifier for type.
Value	text	Generic concept value

The GENCONCEPTTYPES table holds the generic concept types.

Name	Type	Description
Type_Id	long	Primary key for this table. Generic concept type identifier.
Name	text	Name of the generic concept types (e.g. inhibitors, activators, motifs, domains, modifications etc.)



The JOURNAL_SOURCE_REF table holds the journal source information.

Name	Type	Description
Journal_Id	long	Primary key for this table. Journal identifier.
Name	Text	Name of the journal.
Tier	Long	Journal quality tier as classified by New World. (e.g. Tier One, Tier Two, etc.)

The MODIFIESATTRIBUTE table holds the attribute modification information.

Name	Type	Description
Modification_Id	long	Primary key for this table. Attribute modification identifier.
Concept_Id	long	Identifier for concepts.
Operator_Id	long	Identifier for operator.
Attribute_Id	long	Identifier for attribute.
StoredValue	text	Value of attribute modification.
Туре	text	Attribute type

The NAMES table holds the name information.

Name	Type	Description
Name_Id	long	Primary key for this table. Name identifier.
Name	text	Value of the name.
Type_Id	long	Name type

The NAMETYPE table holds the name type definitions.

Name	Туре	Description
Type_Id	long	Primary key for this table. Name type identifier.
Name	text	Value of the name type (e.g. preferred name, synonym, official name, etc.)

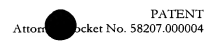
The OPERATOR table holds the operator definitions.

Name	Туре	Description
Operator_Id	long	Primary key for this table. Operator identifier.
Name	text	Value of the operator name (e.g. multiply, set,
		increment, etc.)

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The PACKAGES table holds the client package information.

Name	Type	Description
Package_Id	long	Primary key for this table. Package identifier.
User_Id	long	Identifier for the user.
Name	text	Package name.
Description	text	Package description

The PREPARATIONNAME table holds the information preparations.

Name	Туре	Description
Preparation_Id	long	Primary key for this table. Preparation identifier.
Name	text	The name of the preparation.

The REACTIONS table holds the reaction relationship information.

Name	Type	Description
Reaction_Id	long	Primary key for this table. Reaction identifier.
Event_Id	long	Identifier for events.
Concept_Id	long	Identifier for concepts.
Type_Id	long	Identifier for reaction type.
Stoichiometry_Data	long	Reaction stoichiometry.

The REACTIONTYPE table holds the reaction types.

Name	Type	Description
Type_Id	long	Primary key for this table. Reaction identifier.
Name	text	The name of the reaction. (e.g. requires, inhibited by, produces, excluded from, present in, etc.)

The REAGENTS table holds the event reagent information.

Name	Туре	Description
Concept_Id	. long	Primary key for this table. Concept identifier.
Reference_Id	long	Identifier for references.
Name	long	Reagent name.

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The REFERENCE table holds the reference information.

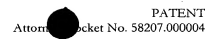
Name	Type	Description
Reference_Id	long	Primary key for this table. Reaction identifier.
Journal	long	Identifier for journal.
Title	text	Reference title.
Year	long	Reference year.
Volume	text	Referene volume.
Issue	text	Reference issue.
Page_Start	text	Reference start page.
Page_End	text	Reference end page.
PMID	long	Reference PMID.
Review	bool	Reference review indicator.

The RELATIONSHIPTYPE table holds the relationship type definitions.

Name	Type	Description
Type_Id	long	Primary key for this table. Relationship type identifier.
Name	long	Relationship type name.

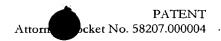
The SECURITY table holds the concept and event security relationships.

Name	Type	Description
Туре	long	Composite key for this table with Event_Id and
		Concept_Id. Security type.
Event_Id	long	Composite key for this table with Type and
		Concept_Id. Event identifier.
Concept_Id	long	Composite key for this table with Type and
		Event_Id. Concept identifier.
Package_Id	long	Security package identifier.



The TESTSATTRIBUTE table holds the attribute information for the tests functionality.

Name	Туре	Description
Test_Id	long	Primary key for this table. Test identifier.
Concept1	long	First concept identifier for the test functionality.
Compare	long	Comparison identifier.
Attribute1	long	First attribute identifier for the test functionality.
Attribute2	long	Second attribute identifier for the test functionality.
Concept2	long	Second concept identifier for the test functionality.



The TIER table holds the reference tier definitions.

Name	Type	Description
Tier_Id	long	Primary key for this table. Tier identifier.
Description	text	The description of the tier. (e.g. Tier One, Tier Two, etc.)

The USERS table holds the user security information information.

Name	Туре	Description
User_Id	long	Primary key for this table. User identifier.
Password	text	User password.
Login	text	Login information for the user.
Full_Name	text	User full name.

The VERSION table holds the user edit trail for data edited in the database.

Name	Type	Description
Version_Id	long	Primary key for this table. User identifier.
Name	text	User password.
VersionDate	text	Login information for the user.
ActionType	text	Edit action type. (e.g. Creation, Editing, Deletion, etc.)

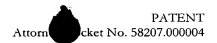
The VERSIONJUNCTION table holds the version information for concepts and events.

Name	Type	Description
Version_Id	long	Primary key for this table. Version identifier.
Concept_Id	long	Concept identifier.
Event_Id	long	Event identifier.

In another embodiment of the present invention, the aforementioned information may be enhanced to model the three dimensional conformation of identified regions (in a protein under study) that might mediate protein-protein interactions. In a further embodiment of the present invention, reference links at key program sites provide the User 20 rapid access to more detailed published information. Hyperlinks are

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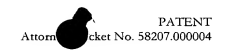


also provided at suitable locations pointing to organism-specific databases.

In another embodiment of the present invention, a User 20 may examine more closely the molecular details of any component in the proposed pathway. This ability has the advantage of allowing the User 20 to effectively zoom in and out on any pathway element for a closer (molecular) or broader (sub-cellular or cellular) look. For example, a a User 20 may take a closer look at the three dimensional conformation of a binding site and its interaction with the target site. The effect of sequence alterations or covalent modification of binding site subunits (e.g., by phosphorylation) may also be examined more closely. This information is advantageous in drug design studies, prediction of toxicity and side effects, and susceptibility issues due to genetic variation among diverse populations.

Although the present invention has been described in terms of particularly preferred embodiments, it is not limited to these embodiments. Alternative embodiments and modifications which would still be encompassed by the invention may be made by those skilled in the art, particularly in light of the foregoing teachings. For example, although the present invention has been described in connection with the simulation of biochemical pathways, it could be used for other applications. For example, the Inference Engine 14 of the Simulation Module 10 may be adapted to process information to predict automotive or computer network traffic. For this type of application, the Database 80 may contain known traffic concepts (e.g., cars, trucks, different types of weather, accidents, etc.) and known traffic events. These concepts and events may then be processed by the Inference Engine 14 of the Simulation Module 10 to predict traffic effects given a traffic stimulus.





Accordingly, this invention is intended to cover any alternative embodiments, modifications or equivalents which may be within the spirit and scope of the invention.